



PCT

**WORLD INTELLECTUAL PROPERTY ORGANIZATION**  
**International Bureau**

INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

***FOR THE PURPOSES OF INFORMATION ONLY***

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	GB	United Kingdom	MR	Mauritania
AU	Australia	GE	Georgia	MW	Malawi
BB	Barbados	GN	Guinea	NE	Niger
BE	Belgium	GR	Greece	NL	Netherlands
BF	Burkina Faso	HU	Hungary	NO	Norway
BG	Bulgaria	IE	Ireland	NZ	New Zealand
BJ	Benin	IT	Italy	PL	Poland
BR	Brazil	JP	Japan	PT	Portugal
BY	Belarus	KE	Kenya	RO	Romania
CA	Canada	KG	Kyrgyzstan	RU	Russian Federation
CF	Central African Republic	KP	Democratic People's Republic of Korea	SD	Sudan
CG	Congo	KR	Republic of Korea	SE	Sweden
CH	Switzerland	KZ	Kazakhstan	SI	Slovenia
CI	Côte d'Ivoire	LI	Liechtenstein	SK	Slovakia
CM	Cameroon	LK	Sri Lanka	SN	Senegal
CN	China	LU	Luxembourg	TD	Chad
CS	Czechoslovakia	LV	Latvia	TG	Togo
CZ	Czech Republic	MC	Monaco	TJ	Tajikistan
DE	Germany	MD	Republic of Moldova	TT	Trinidad and Tobago
DK	Denmark	MG	Madagascar	UA	Ukraine
ES	Spain	ML	Mali	US	United States of America
FI	Finland	MN	Mongolia	UZ	Uzbekistan
FR	France			VN	Viet Nam
GA	Gabon				

TITLE OF INVENTION  
HAEMOPHILUS OUTER MEMBRANE PROTEIN

FIELD OF INVENTION

5       The present invention is related to the field of molecular genetics and is particularly concerned with the cloning of an outer membrane protein D15 of Haemophilus.

BACKGROUND OF THE INVENTION

10      Haemophilus influenzae type b (Hib) is a major cause of bacterial meningitis in children under the age of five years. Protective antibodies to the disease are induced by the capsular polysaccharide of the organism and a vaccine was developed that utilises the purified polyribosyl ribitol phosphate (PRP) as the antigen. This 15 vaccine provides 90% protection in adults and in children over 24 months of age, but was ineffective in children under 24 months Zangwill et al 1993 (The references are identified in a list of reference at the end of this disclosure). Like other polysaccharide antigens, PRP 20 does not induce the proliferation of T-helper cells, and re-immunisation fails to elicit either a booster response or an increase in memory cells. Conjugation of the PRP polysaccharide with protein carriers confers T-cell dependent characteristics to the vaccine and 25 substantially enhances the immunologic response to the PRP antigen. Currently, there are four PRP-carrier conjugate vaccines available. These are vaccines based upon H. influenzae type b capsular polysaccharide conjugated to diphtheria toxoid, tetanus toxoid, or 30 Neisseria meningitidis outer membrane protein (reviewed in Zangwill et al 1993).

However, the current Haemophilus conjugate vaccines only protect against meningitis caused by Haemophilus influenzae type b. They do not protect against other 35 invasive typeable strains (types a and c) and, more importantly, against non-typeable (NTHi) strains which are a common cause of postpartum and neonatal sepsis,

pneumonia and otitis media. In the United States alone, treatment of otitis media costs between 1 and 2 billion dollars per year for antibiotics and surgical procedures, such as tonsillectomies, adenoidectomies and insertion of 5 tympanostomy tubes. To achieve universal protection against H. influenzae related diseases in the 2 to 6 month age group and certain high risk groups, the provision of conserved, cross-reactive non-capsular H. influenzae immunogens is desirable. Methods for 10 inducing immunity against disease are constantly improving and there is presently a move to use subunits and better defined materials as antigens. This is being undertaken to minimise or eliminate potential side-effects caused by certain native immunogens, while 15 preserving their immunogenicity to confer protection against the disease. Therefore, it would be very attractive to develop a universal vaccine against Haemophilus using cross-reactive outer membrane proteins, fragment, analogs, and/or peptides corresponding thereto 20 as protective antigens. Such antigens may be incorporated into the conventional H. influenzae type b conjugate vaccines as additional immunogens or used as autologous carriers for H. influenzae capsular polysaccharides. A high molecular weight outer membrane 25 protein D15 found in non-typeable and type b stains of H. influenzae has been identified as a cross-reactive antigen (Thomas et al., 1990). D15 appears to be cell surface-exposed in its natural state and exhibits a molecular mass of about 80 kDa as judged by SDS-PAGE 30 analysis. It would be desirable to provide the sequence of the DNA molecule that encodes this D15 outer membrane protein and peptides corresponding to portions thereof for diagnosis, immunization and the generation of diagnostic and immunological reagents. The diseases 35 caused by Haemophilus are serious and improved methods for preventing, detecting and treating diseases such as

otitis media, epiglottitis, pneumonia, and tracheobronchitis, are required.

#### SUMMARY OF THE INVENTION

The present invention is directed towards the provision of purified and isolated nucleic acid molecules comprising at least a portion coding for a D15 outer membrane protein of a species of Haemophilus. The nucleic acid molecules comprising at least a portion coding for D15 outer membrane protein are useful for the specific detection of strains of Haemophilus, and for diagnosis of infection by Haemophilus. The purified and isolated nucleic acid molecules, such as DNA comprising at least a portion coding for D15 outer membrane protein, are also useful for expression of the D15 gene by recombinant DNA means for providing, in an economical manner, purified and isolated D15 outer membrane protein.

The D15 outer membrane protein or fragments thereof or analogs thereof are useful immunogenic compositions for the preparation of vaccines against diseases caused by Haemophilus, the diagnosis of infection by Haemophilus and as tools for the generation of immunological reagents. Mono- or polyclonal antisera (antibodies) raised against the D15 outer membrane protein produced in accordance with aspects of the present invention are useful for the diagnosis of infection by Haemophilus, specific detection of Haemophilus (in, for example, in vitro and in vivo assays) and for the treatment of diseases caused by infection by Haemophilus.

Peptides corresponding to portions of the D15 outer membrane protein or analogs thereof are useful immunogenic compositions for the preparation of vaccines against disease caused by Haemophilus, the diagnosis of infection by Haemophilus and as tools for the generation of immunological reagents. Mono- or polyclonal antisera raised against these peptides, produced in accordance with aspects of the present invention, are useful for the

diagnosis of infection by Haemophilus, specific detection of Haemophilus (in, for example, in vitro and in vivo assays) and for use in passive immunization as a treatment of disease caused by infection by Haemophilus.

5 In accordance with one aspect of the present invention, therefore, there is provided a purified and isolated nucleic acid molecule, the molecule comprising at least a portion coding for a D15 outer membrane protein. The nucleic acid molecule has a DNA sequence  
10 selected from:

(a) the DNA sequence set out in any one of Figures 1A to 1E (as described below) or its complementary strand; and

15 (b) DNA sequences which hybridize under stringent conditions to the DNA sequences defined in (a). The DNA sequences defined in (b) preferably has at least 90% sequence identity with the sequences defined in (a). The DNA sequence defined in (b) particularly may comprise the consensus sequence set forth in Figure 1F (as described  
20 below).

In another aspect of the present invention, there is provided a purified and isolated D15 outer membrane protein or a portion thereof. The D15 outer membrane protein may be a Haemophilus D15 outer membrane protein and more particularly an H. influenzae D15 outer membrane protein and the H. influenzae strain may be an H. influenzae type b strain, such as H. influenzae type b strains Ca or Eagan or MinnA or a non-typeable H. influenzae strain, such as PAK 12085 or SB33.

25 30 35 In an additional embodiment, the present invention also includes a recombinant plasmid adapted for transformation of a host, the recombinant plasmid comprising a plasmid vector into which has been inserted a DNA segment comprising the purified and isolated DNA molecule provided herein. Such recombinant plasmid comprises a plasmid vector into which a DNA segment which

comprises at least an 18 bp fragment selected from the DNA molecules as recited above is inserted. The recombinant plasmid may be plasmid DS-712-2-1 having ATCC accession number 75604, deposited November 4, 1993 and 5 plasmid JB-1042-5-1 having ATCC accession number 75006, deposited November 4, 1993.

The plasmids may be adapted for expression of the encoded D15 outer membrane protein in a host cell, which may be a heterologous or homologous host, by 10 incorporation into a recombinant vector, provided in accordance with a further aspect of the invention. The recombinant vector may comprise at least a DNA segment comprising at least an 18 bp fragment selected from the DNA molecules as recited above and expression means 15 operatively coupled to the DNA segment for expression of the gene product encoded thereby in the host cell. The plasmid for expression of the encoded D15 outer membrane protein may be plasmid DS-880-1-2 having ATCC accession number 75605, deposited November 4, 1993 being adapted 20 for expression at the D15 outer membrane protein in E. coli. The selected DNA segment may encode a polypeptide of at least 6 residues and, in particular, may be selected from those segments encoding a polypeptide of 25 Table 2 (below). The DNA segment may further comprise a nucleic acid sequence encoding a leader sequence for export of the gene product from the host. The host for expression may be selected from, for example, Escherichia coli, Bacillus, Haemophilus, fungi, yeast or the baculovirus expression system may be used.

30 Additional aspects of the invention include the protein encoded by the DNA molecule comprising at least a portion coding for the D15 outer membrane protein, fragment or a functional analog of such protein, the use 35 of the protein or analog in vaccination and diagnosis, and the generation of immunological reagents. The invention also includes antisera (antibodies) raised

against the D15 outer membrane protein encoded by the DNA molecule comprising at least a portion coding for a D15 outer membrane protein and purified peptides corresponding to portions of the D15 outer membrane  
5 protein and there are in passive immunization and treatment of diseases caused by Haemophilus.

According to another aspect of the invention, a purified and isolated peptide containing an amino acid sequence corresponding to the amino acid sequence of at  
10 least a portion of the D15 outer membrane protein or variant or mutant which retains immunogenicity. The peptide may be produced by recombinant methods or peptide synthesis whereby the purified peptide is free from contaminants associated with bacteria normally containing  
15 the D15 outer membrane protein. Such synthetic peptides preferably have an amino acid sequence selected from those presented in Table 2.

In accordance with an additional aspect of the invention, an immunogenic composition is provided which  
20 comprises the D15 outer membrane protein, fragments thereof, functional analogs thereof, or peptides as recited above and a physiologically-acceptable carrier therefor. Such immunogenic composition is particularly formulated as a vaccine for in vivo administration to  
25 protect against diseases caused by Haemophilus. For such purpose, the immunogenic composition may be formulated as a microparticle preparation, capsule preparation or liposome preparation. In addition, such immunogenic composition may be provided in combination with a  
30 targeting molecule for delivery to specific cells of the immune system or to mucosal surfaces.

In accordance with a further aspect of the invention, there is provided a method for inducing protection against disease caused by Haemophilus,  
35 comprising the step of administering to a subject, including a mammal, such as a human, an effective amount

of the immunogenic composition or the nucleic acid molecule as recited above to provide protective immunity against Haemophilus infection.

The present invention further includes a chimeric 5 molecule comprising a D15 protein or peptide corresponding thereto as provided herein linked to another polypeptide or protein or a polysaccharide. The linked polypeptide or protein may comprise a surface protein or peptide corresponding thereto from a 10 pathogenic bacteria, which may be the P1, P2 or P6 outer membrane protein of H. influenzae. The linked polysaccharide preferably comprise a PRP molecule from H. influenzae.

#### BRIEF DESCRIPTION OF THE FIGURES

15 The present invention will be further understood from the following description with reference to the drawings, in which:

Figure 1A shows the nucleotide sequence of the D15 gene from H. influenzae type b Ca strain (SEQ ID NO: 1) 20 and its deduced amino acid sequence (SEQ ID NO: 2);

Figure 1B shows the nucleotide sequence of the D15 gene from H. influenzae type b Eagan strain (SEQ ID NO. 3) and its deduced amino acid sequence (SEQ ID NO: 4);

Figure 1C shows the nucleotide sequence of the D15 gene from H. influenzae type b MinnA strain (SEQ ID NO. 25 5) and its deduced amino acid sequence (SEQ ID NO: 6);

Figure 1D shows the nucleotide sequence of the D15 gene from H. influenzae non-typeable SB33 (SEQ ID NO. 7) and its deduced amino acid sequence (SEQ ID NO: 8);

30 Figure 1E shows the nucleotide sequence of the D15 gene from H. influenzae non-typeable PAK 12085 (SEQ ID NO. 9) and its deduced amino acid sequence (SEQ ID NO: 10);

Figure 1F shows an alignment of the nucleotide 35 sequences of the D15 genes (SEQ ID NOS: 1, 3, 5, 7 and 9)

obtained from different H. influenzae isolates (typeable, Ca, Eagan and MinnA; nontypeable SB33 and PAK 12085);

Figure 2 shows restriction maps of clones pUC19/D15 (Ca), DS-712-2-1 (Eagan), DS-691-1-5 (MinnA), JB-1042-5-1 5 (SB33), and JB-1042-9-4 (PAK 12085). H = HindIII; R = EcoRI; S = Sau3A I; and Xb = XbaI;

Figure 3 shows an alignment of the amino acid sequences of D15 outer membrane proteins (SEQ ID NOS: 2, 10 4, 6, 8 and 10) obtained from different H. influenzae isolates (typeable, Ca, Eagan and MinnA; nontypeable, SB33 and PAK 12085). Amino acids are represented by the conventional one-letter code. The Ca D15 sequence is used as reference and the dots indicate amino acid residues which are identical to those of the Ca D15 outer 15 membrane protein;

Figure 4 shows the construction of a plasmid (DS-880-1-2) expressing full-length SB33 D15 (rD15) from the strong inducible T7 promoter;

Figure 5 shows an SDS-PAGE analysis of native D15 20 affinity-purified from H. influenzae strain 30;

Figure 6 shows an SDS-PAGE analysis of sequential fractions obtained during the purification of the full-length rD15 expressed in E. coli containing plasmid DS-880-1-2;

Figure 7 shows guinea pig IgG antibody responses to 25 full length rD15. The arrows indicate the immunization schedule. Bleeds were taken at 0, 2, 4, 6 and 8 weeks. The bars represent the standard deviation;

Figure 8 shows mouse IgG antibody responses to full 30 length rD15. The arrows indicate the immunization schedule. Bleeds were taken at 0, 1, 4, 5 and 7 weeks. The bars represent the standard deviation;

Figure 9 shows an SDS-PAGE analysis of the N-terminal rD15 fragment purified from GST-(D15 fragment) 35 fusion protein. Lanes: 1, prestained low molecular weight markers (14kDa, 21kDa, 31 kDa, 45kDa, 68kDa,

97kDa); 2, GST standard; 3, GST-(D15 fragment) fusion protein; 4, fusion protein cleaved by thrombin; 5, N-terminal rD15 fragment; 6, GST; 7, low molecular weight markers;

5       Figure 10 shows guinea pig IgG antibody response to N-terminal rD15 fragment. The arrows indicate the immunization schedule. Bleeds were taken at 2, 4, 6 and 8 weeks. The bars represent the standard deviation;

10      Figure 11 shows the hydrophilicity plot of D15 established by using a window average across 7 residues according to Hope, 1986.

#### GENERAL DESCRIPTION OF THE INVENTION

Any Haemophilus strains that have D15 genes may be conveniently used to provide the purified and isolated nucleic acid molecules (which may be in the form of DNA molecules), comprising at least a portion coding for a D15 outer membrane protein as typified by embodiments of the present invention. Such strains are generally available from clinical sources and from bacterial culture collections, such as the American Type Culture Collection. H. influenzae strains may include types a, b and c strains, non-typeable strains and other bacteria that produce a D15 protein, fragment or analog thereof.

Appropriate strains of Haemophilus include:-

25      H. influenzae type b strain Ca;  
          H. influenzae type b strain MinnA;  
          H. influenzae type b strain Egan;  
          H. influenzae non-typeable b strain SB33; or  
          H. influenzae non-typeable b strain PAK 12085.

30      In this application, the term D15 outer membrane protein is used to define a family of D15 proteins which includes those having naturally occurring variations in their amino acid sequences as found in various strains of, for example, Haemophilus. The purified and isolated DNA molecules comprising at least a portion coding for D15 outer membrane protein of the present invention also

include those having naturally occurring variations in their nucleic acid sequences as found in various strains of, for example Haemophilus and those DNA molecules encoding functional analogs of D15 outer membrane protein. In this application, a first protein is a functional analog of a second protein if the first protein is immunologically related with and/or has the same function as the second protein. The functional analog may be, for example, a fragment of the protein or 10 a substitution, addition or deletion mutant thereof.

In aspects of the present invention, the D15 gene was isolated from H. influenzae type b strain Ca as shown in Figure 1A; H. influenzae type B Eagan, Figure 1B; H. influenzae type b MinnA, Figure 1C; non-typeable H. influenzae SB33, Figure 1D; non-typeable H. influenzae PAK 12085, Figure 1E. A comparison of the nucleic acid sequences of the D15 genes and of the deduced amino acid sequences of the D15 outer membrane proteins from these strains of H. influenzae showed the genes and proteins to 20 be highly conserved (Figures 1F and 3). The consensus sequence (SEQ ID NO: 55) for the D15 gene is shown in Figure 1F.

The purified and isolated DNA molecules comprising at least a portion coding for a D15 outer membrane protein of a species of Haemophilus, typified by the embodiments described herein, are advantageous as:

- nucleic acid probes for the specific identification of Haemophilus strains in vitro or in vivo;
- 30 - the products encoded by the DNA molecules are useful as diagnostic reagents, antigens for the production of Haemophilus-specific antisera, for vaccination against the diseases caused by species of Haemophilus and detecting infection by Haemophilus; and
- 35 - peptides corresponding to portions of the D15 outer membrane protein as typified by the embodiments

described herein are advantageous as diagnostic reagents, antigens for the production of Haemophilus-specific antisera, for vaccination against the diseases caused by species of Haemophilus and for detecting infection by 5 Haemophilus.

Reference will now be made in detail to the presently preferred embodiments of the invention, which together with the following Examples, serve to explain the principle of the invention. For clarity of 10 disclosure, and not by way of limitation, the detailed description of the invention is divided into the following sections:

(i) The DNA sequences coding for the outer membrane protein D15 from H. influenzae type b Ca strain.

A clone producing the outer membrane protein designated D15 of H. influenzae type b (Hib) was isolated by screening a genomic library with H. influenzae type b OMP-specific polyclonal antibodies as previously described by Berns and Thomas 1965; Thomas and Rossi 20 1986. The DNA fragment encoding the D15 protein was isolated, subcloned into pUC19 to produce pUC19/D15 (Figure 2) and used to transform E. coli HB101 as described in Example 1. Plasmid DNA was prepared from two individual colonies of E. coli HB101 containing the 25 pUC19/D15 plasmid. Sequencing was performed on an ABI DNA sequencer model 370A using dye-terminator chemistry and oligonucleotide primers which had been synthesized on an ABI DNA synthesizer model 380B, and purified by chromatography. Nucleotide sequence analysis of the D15 30 gene revealed that it contains a putative promoter and an open reading frame encoding 789 amino acids (Figure 1A).

The first 19 amino acid residues of the translated open reading frame form a typical leader sequence as found in other H. influenzae type b outer membrane 35 proteins, such as P1 and P2. The N-terminal sequence of immuno-affinity purified native D15 antigen was

determined by automated Edman degradation using the ABI 477A protein sequencer and was found to be Ala-Pro-Phe, which is identical to the N-terminal amino acid sequence Ala-Pro-Phe-Val-Ala-Lys- (SEQ ID NO: 11) predicted from 5 an analysis of the sequence of the D15 gene presented in Figure 1A.

(ii) **The sequence of D15 genes from other H. influenzae strains.**

D15 genes were isolated from other H. influenzae strains by screening the chromosomal libraries of H. influenzae type b strains Eagan, Minn A and the non-typeable H. influenzae (NTHi) strains SB33 and PAK 12085, as described in Examples 2, 3 and 4. Hybridization-positive clones were plated and submitted to a second 15 round of screening. The restriction maps of the clones obtained are shown in Figure 2. The nucleotide sequences of the D15 genes were determined for all these clones (Figures 1B to 1E) and their derived amino acid sequences compared (Figure 3). The D15 amino acid 20 sequences of the three H. influenzae type b strains were identical and only a few amino acid differences were observed in the amino acid sequence of the D15 protein from the non-typeable strains (Figure 3).

(iii) **Expression of D15 and its fragments in E. coli.**

25 Since D15 is expressed in small quantities by strains of H. influenzae, it is advantageous to either express this antigen as a recombinant protein in a heterologous system, such as E. coli, or to modify the H. influenzae organism to enhance native D15 expression. The 30 Hind III/Eco RI fragment of H. influenzae type b Ca strain DNA encoding the full length D15 protein was expressed in pUC19 but not pUC18, suggesting that the lac promoter is helping to express the D15 gene in E. coli, even though the native D15 gene promoter is present. The 35 T7 expression system is a tightly controlled, inducible system which has great utility in expression of

heterologous proteins in E. coli. The T7 expression system is described in U.S. Patent 4,952,496. Clones were, therefore, constructed which utilize the T7 system to express a mature D15 protein that contains an additional methionine residue at the amino terminus. The D15 signal sequence was removed during this construction process. A full length recombinant D15 (termed rD15) was expressed in inclusion bodies which allow the D15 protein to be readily purified. The D15 genes from H. influenzae type b strain Ca and H. influenzae non-typeable SB33 strain have been expressed at high levels in E. coli using the T7 system to permit production of large quantities of rD15 protein. The construction of clone DS-880-1-2 which expresses the SB33 D15 gene is described herein (see Figure 4 and Example 5). The rD15 protein was immunologically similar to its native counterpart isolated from H. influenzae typeable and non-typeable strains (see below). Thus, rD15 may be used as a cross-reactive antigen in a diagnostic kit to detect many, if not all, strains of H. influenzae and other bacteria that produce a D15 outer membrane protein or analog thereof. Alternatively, rD15 can be used as an antigen to specifically detect the presence of H. influenzae in a sample.

A truncated D15 fragment was expressed in E. coli as a fusion protein with glutathione S-transferase (GST), as described in Example 6. The construction was designed to express the N-terminal fragment of the D15 protein. The fusion protein was expressed at high levels from a pGEX-2T construction and the N-terminal fragment was cleaved from the GST carrier protein by treatment with thrombin. This procedure generated a molecule termed the N-terminal rD15 fragment which encompasses amino acids 63-223 of the D15 protein. This N-terminal rD15 fragment was highly immunogenic and elicited protective antibodies against challenge with live H. influenzae.

(iv) Purification of native D15 from H. influenzae cell paste.

The present invention also provides a method to prepare purified native D15 protein from H. influenzae.

- 5 The protein is extracted and affinity-purified from the cell pastes of either H. influenzae typeable or non-typeable isolates by a procedure involving the dissolution of the protein in an aqueous detergent solution (see Example 13). The native D15 protein from a  
10 non-typeable H. influenzae strain 30 was solubilized with a 50 mM Tris-HCl/ 0.5% Triton X-100/ 10 mM EDTA buffer, pH 8.0 and further purified on a D15-specific monoclonal antibody affinity column (Figure 5A). An 80 kDa protein was eluted from the column with 50 mM diethylamine, pH  
15 12.0 and shown to react with a D15-specific monoclonal antibody on immunoblot analysis (Fig. 5B). The native D15 is also highly immunogenic in experimental animals. Rabbit anti-D15 antisera reacted with all H. influenzae isolates as determined by immunoblot analyses.  
20 (v) Purification of a full-length recombinant D15 protein expressed in E. coli.

A full-length recombinant D15 (rD15) protein was expressed in inclusion bodies in E. coli. As shown in Figure 6, purification of rD15 inclusion bodies was achieved by a sequential extraction of the E. coli cell lysate with 50 mM Tris-HCl, pH 8.0, then 50 mM Tris containing 0.5% Triton X-100 and 10 mM EDTA, pH 8.0. After centrifugation, more than 95% of the proteins in the resulting pellet was an 80 kDa protein by SDS-PAGE analysis, that reacted with a D15-specific monoclonal antibody on an immunoblot. The N-terminal sequence of the rD15 was found to be Met-Ala-Pro-Phe-Val-Lys-Asp- (SEQ ID NO: 54) which is identical to the predicted amino acid sequence.

- 35 The rD15 inclusion bodies were solubilized with a mixture of PBS, 0.5% Triton X-100, 10 mM EDTA and 8 M

urea (see Example 8). After dialysis against PBS to remove urea, more than 80% of the D15 protein remained soluble. This soluble rD15 antigen was used for the immunogenicity studies described below. From shake-flask experiments, it was estimated that about 10 mg of soluble rD15 protein was obtained from 1 L of E. coli bacterial culture. It is clear that growing the recombinant E. coli strains under optimised fermentation conditions significantly increase the level of rD15 production.

10 (vi) Immunogenicity of the full-length recombinant D15 protein (rD15).

The immunogenicity of the full-length rD15 protein was studied in guinea pigs and mice. Using the immunization protocols described in Figure 7, a 15 µg dose of rD15 induced high IgG titers in guinea pigs when administered in the presence of either Freund's adjuvant or AlPO<sub>4</sub>. In the mouse dose-response study, the protein appeared to be immunogenic at a dose as low as 5 µg in either Freund's adjuvant (Figure 8A) or AlPO<sub>4</sub> (Figure 20 8B).

The protective ability of rD15 against H. influenzae type b infection was examined in the infant rat model of bacteremia essentially as described by Loeb (1987). Thus, infant rats passively immunized with guinea pig anti-rD15 antisera were significantly less bacteremic than controls injected with pre-bleed sera, which is consistent with the previous report by Thomas et al. (1990).

25 (vii) Purification and characterization of the N-terminal rD15 fragment.

The truncated rD15 fragment corresponding to the N-terminus of the D15 protein (residues 22 to 223) as described in Example 6, was expressed in E. coli as a soluble protein fused to GST. The fusion protein (46 kDa) was readily extracted using phosphate buffered saline (PBS). Purification of the GST-D15 fragment fusion

protein was achieved by a single-step affinity purification process on a glutathione-Sepharose 4B column (Figure 9, Lane 3). Cleavage of the 46 kDa fusion protein with thrombin yielded two fragments (Figure 9, 5 Lane 4), a 26 kDa protein which corresponded to a purified GST standard (Figure 9, Lane 2), and a 20 kDa polypeptide which had the size expected for the N-terminal rD15 fragment (amino acid residues 63 to 223), respectively. Separation of these two proteins was 10 achieved by a second round of glutathione-Sepharose 4B affinity chromatography. From shake-flask experiments, it was estimated that about 1 mg of purified N-terminal rD15 fragment was recovered from 1 L of E. coli bacterial culture. It is clear that growing the recombinant E. 15 coli strains under optimised fermentation conditions will significantly increase the level of N-terminal rD15 fragment production.

The identity of the 20 kDa polypeptide and the 26 kDa protein was confirmed by both immunoblotting and 20 protein sequencing. The N-terminal sequence of the 20 kDa polypeptide was found to be NH<sub>2</sub>-Ser-Leu-Phe-Val-Ser-Gly-Arg-Phe-Asp-Asp-Val-Lys-Ala-His-Gln-Glu-Gly-Asp-Val-Leu-Val-Val-Ser- (SEQ ID NO: 12), which corresponds to residues 63 to 85 of the primary sequence of D15. This 25 result indicates that there is a spurious thrombin cleavage site within the D15 sequence and that the first 42 amino acids of the rD15 fragment are cleaved off during thrombin digestion. Thus, the final N-terminal rD15 fragment was 161 amino acids in length corresponding 30 to residues 63 to 223 of the primary sequence of D15. The N-terminal sequence obtained for the 26 kDa protein (NH<sub>2</sub>-Met-Ser-Pro-Ile-Leu-Gly-Tyr-Trp-Lys- - SEQ ID NO: 13) confirmed that it was GST.

(viii) Immunogenicity of the N-terminal rD15 fragment.  
35 The immunogenicity of the N-terminal rD15 fragment was tested in guinea pigs using various adjuvants. Using

the immunization protocols described in Figure 10, a 10 µg dose of N-terminal rD15 fragment induced a good booster response in guinea pigs with almost all the adjuvants tested. The highest anti-D15 IgG titer was 5 observed in the group of guinea pigs immunized with N-terminal rD15 fragment in Freund's adjuvant. The second best adjuvant was Titermax (CytRx Inc.). The other two adjuvants, TPAD4 (tripalmitoyl-Cys-Ser-Glu<sub>4</sub>) and AlPO<sub>4</sub> were equally potent.

10 (ix) **Protective ability of the N-terminal rD15 fragment against H. influenzae type b challenge.**

An in vivo challenge model for assessing the protective abilities of antigen against diseases caused by Haemophilus is the infant rat model of bacteremia as 15 described by Loeb 1987. The protective ability of the N-terminal rD15 fragment against H. influenzae type b challenge was examined in this rat model. As illustrated in Table 1, infant rats passively immunized with rabbit 20 anti-N-terminal rD15 fragment antisera showed significantly lower bacteremia compared to those injected with pre-bleed sera.

Since passively transferred antisera against the N-terminal rD15 fragment were found to be protective in the infant rat model of bacteremia, it was of interest to 25 identify the protective epitope(s) of this N-terminal rD15 fragment. The first nine overlapping peptides of the D15 protein as listed in Table 2 were chemically synthesized based upon the amino acid sequence derived from the sequence of the D15 gene from H. influenzae type 30 b Ca (Figure 1). These synthetic peptides were assessed for their reactivities with either rabbit or guinea pig antisera raised against purified N-terminal rD15 fragment by ELISAs. As shown in Table 3, both guinea pig and rabbit antisera reacted with a cluster of D15 peptides, 35 including peptides D15-P4 to D15-P8 encompassing residues 93 to 209 of the D15 primary sequence.

Further studies were performed to determine whether the protection against H. influenzae type b observed using rabbit anti-D15 antisera in infant rats could be neutralized by D15 peptides. In the first experiment, a 5 rabbit anti-N-terminal rD15 fragment antiserum was injected into a group of seven infant rats in the presence or absence of a mixture of the nine D15 peptides (D15-P2 to D15-P10). Animals in the positive control group were injected with the rabbit anti-N-terminal rD15 10 fragment antiserum mixed with purified D15 fragment and the negative control group was injected with a mixture of the nine peptides only. As illustrated in Table 4, infant rats passively immunized with a rabbit anti-N-terminal rD15 fragment antiserum (group #1) showed a significantly 15 lower bacteremia level (3%,  $p = 1.2 \times 10^{-7}$ ) compared to those in the negative control group (group #4, 100%), which was consistent with the previously obtained results. The protection mediated by the rabbit anti-N-terminal rD15 fragment antiserum was largely neutralized 20 by the addition of purified N-terminal rD15 fragment (group #3, 64%), as indicated by the lack of significant difference in the bacteremia level between group #3 and group #4 ( $p = 0.09$ ). Although the addition of the mixture 25 of nine D15 peptides only slightly neutralized the protection conferred by the antiserum (group #2, 13%) as compared to group #1 (3%), the difference in bacteria counts between these two groups was statistically significant ( $p = 0.0037$ ).

To more clearly define the protective epitope(s) of 30 the N-terminal rD15 fragment, the above experiment was repeated with a mixture of five peptides (peptides D15-P4 to D15-P8) which were chosen for their strong reactivities with the rabbit anti-N-terminal rD15 fragment antiserum. The results obtained from this second 35 experiment showed that the protection observed using rabbit anti-N-terminal rD15 fragment (Table 5, group #1)

was completely blocked by the addition of this mixture of five peptides (Table 5, group #2, 106%,  $p = 0.53 \times 10^{-8}$ ). These results strongly indicate that a cocktail of D15 synthetic peptides may be used as immunogens to induce protective antibodies against H. influenzae.

5 (x) **Epitope prediction and peptide synthesis.**

To map the immunodominant T-cell or B-cell epitopes of D15, overlapping synthetic peptides covering the entire D15 protein sequence (Table 2 - SEQ ID NO: 14 to 10 49) were synthesized using the t-Boc solid-phase peptide synthesis as described in Example 15. The peptides were chosen based on their high index of hydrophilic  $\beta$ -turns estimated by secondary structure prediction analysis (Figure 11). Such peptides are likely to be surface-exposed and antigenic. Peptides more than 25 residues in 15 length were selected to better mimic native epitopes.

(xi) **Identification and characterization of immunodominant epitopes of D15 using synthetic peptides.**

To map the linear B-cell epitopes of D15, 20 overlapping synthetic peptides representing the entire sequence of D15 were individually coated onto ELISA plates and probed with several anti-rD15 antisera as described in Example 19. The results are summarized in Table 6. Mouse antisera raised against rD15 reacted with 25 all D15 peptides, but the major epitopes were located within peptides D15-P8 (residues 180-209 - SEQ ID NO: 21), D15-P10 (residues 219-249 - SEQ ID NO: 23), D15-P11 (residues 241-270 - SEQ ID NO: 24), and D15-P26 (residues 554-582 - SEQ ID NO: 39), respectively. Rabbit anti-D15 30 antisera recognized only peptides D15-P4 (residues 93-122 - SEQ ID NO: 17), D15-P14 (residues 304-333 - SEQ ID NO: 27) and D15-P36 (residues 769-798 - SEQ ID NO: 49). Guinea pig antisera raised against rD15 reacted with peptides D15-P2 (residues 45-72 - SEQ ID NO: 15), D15-P4 (residues 93-122 - SEQ ID NO: 17), D15-P6 (residues 135- 35 164 - SEQ ID NO: 19), D15-P8 (residues 180-209 - SEQ ID

NO: 21), D15-P14 (residues 304-333 - SEQ ID NO: 27), D15-P27 (residues 577-602 - SEQ ID NO: 40). The immunodominant linear B-cell epitopes of D15 were thus found to be located within peptides D15-P4 (residues 93-122 - SEQ ID NO: 17) and D15-P14 (residues 304-333 - SEQ ID NO: 27), since these are the only two peptides recognized by rD15-specific antisera from all three animal species. These results indicate that the peptides containing the linear B-cell epitope sequences described above can be used as target antigens in, for example, diagnostic kits to detect the presence of anti-D15 and anti-H. influenzae antibodies in samples.

**(xii) Identification and characterization of immunodominant T-cell epitopes of D15 using synthetic peptides.**

The importance of cytokine networks in the immune and inflammatory responses in immunity and inflammation and their alteration in pathology is becoming more evident as new members of the cytokine family are identified and characterized. Mills et al. (1993) have recently reported that there is a rapid clearance of B. pertussis from the lungs of mice on challenge six weeks after respiratory infection or following two immunizations with the whole-cell pertussis vaccine. Spleen cells from these immunized mice were found to secrete high levels of IL-2 and IFN- $\gamma$  and low levels of IL-5 in the presence of pertussis antigen (pertussis toxoid, filamentous haemagglutinin (FHA) and pertactin). This result suggests that Th1 cell (T-cells producing high levels of IL-2 and IFN- $\gamma$ ) proliferation is very important for recovering from respiratory infection. The generation of Th1 and Th2 cell subsets is regulated by the balance between different groups of cytokines, predominantly IL-12 and IL-4 (Trinchieri, 1993). IL-12 and IL-4 are responsible for Th1 and Th2 cells differentiation, respectively. One of the roles of Th2

cells in the immune system is to provide helper activity for eliciting high levels of antigen-specific antibodies following immunization. Antigens containing Th1 epitope(s) stimulate antigen-specific T-cells to produce 5 high levels of IL-2 and IFN- $\gamma$ , whereas Th2 epitope(s) induce high levels of IL-4 expression. Th0 epitope(s) stimulate the synthesis of IFN- $\gamma$  and IL-4.

Little is known about the cellular immune response to outer membrane proteins of H. influenzae and its role 10 in the protection against H. influenzae infection and diseases. To this end, the inventors performed studies of the cellular response elicited in mice following rD15 immunization. D15-specific T-cell epitopes were determined using D15 peptides and T-cell lines obtained 15 from five BALB/c mice immunized with rD15 (see Example 23). The lymphocyte proliferative responses of the D15-specific T-cell lines to overlapping D15 peptides were determined in conventional cytokine assays as described in Example 24. The results summarized in Table 7, 20 revealed that stimulation only with certain synthetic peptides elicited proliferative responses and the release of specific cytokines. Synthetic peptides corresponding to residues 114-143 (D15-P5 - SEQ ID NO: 18), 282-312 (D15-P13 - SEQ ID NO: 26) and 577-602 (D15-P27 - SEQ ID 25 NO: 40), and 219-249 (D15-P10 - SEQ ID NO: 23), 262-291 (D15-P12 - SEQ ID NO: 25), 390-416 (D15-P18 - SEQ ID NO: 31), 410-435 (D15-P19 - SEQ ID NO: 32) 554-582 (D15-P26 - SEQ ID NO: 39), 596-625 (D15-P28 - SEQ ID NO: 41), 725- 750 (D15-P34 - SEQ ID NO: 47) and 745-771 (D15-P35 - SEQ 30 ID NO: 48) were shown to be highly stimulatory for rD15-specific BALB/c Th0 cells and Th1 cells, respectively. Therefore, these immunodominant T-cell epitopes can be used as autologous carriers for PRP, and/or OMP B-cell epitopes to enhance their immunogenicity. The Th1 cell 35 epitopes identified above may be useful in the H.

influenzae vaccine formulations to induce H. influenzae-specific cellular immune responses.

(xiii) **Immunogenicity of D15 peptides.**

To determine whether synthetic D15 peptides were immunogenic free peptides were assessed individually for their immunogenicity. Rabbit and guinea pig anti-peptide antisera were tested for their reactivities with the immunizing peptides as well as with native D15 and rD15 by ELISA and immunoblotting. As shown in Table 8, all guinea pig anti-D15 peptide antisera except those raised against D15-P26 (SEQ ID NO: 39), D15-P29 (SEQ ID NO: 42), D15-P30 (SEQ ID NO: 43) and D15-P31 (SEQ ID NO: 44) were shown to be immunogenic by ELISAs. The induction of high titers of peptide-specific IgG antibodies by free peptides clearly indicates that most peptides contain both a functional T-helper determinant and a B-cell epitope(s). In addition, these anti-peptide antisera recognised D15 in the immunoblot assay. Since most peptides contain potent functional T-helper determinant(s) and induce strong IgG antibody responses in mammals, they are candidate immunogens for inclusion in an H. influenzae vaccine preparation. D15 peptide-specific antisera cross-reacted with D15 from non-typeable strains of H. influenzae as judged by immunoblotting. This finding indicates that immunogenic D15 peptides contain epitopes which are highly conserved among typeable and non-typeable strains of H. influenzae. In addition, polyclonal antibodies against these epitopes are useful to detect H. influenzae in biological samples.

Therefore, these conserved epitopes of D15 can be used either individually or in combination to prepare cross-reactive synthetic immunogens against typeable and non-typeable strains of H. influenzae and other bacteria that produce D15 protein, a fragment or an analog thereof. Peptides described above can be further polymerized, or modified with lipids as lipopeptides, or

linked to polysaccharides including PRP as synthetic glycopeptide or lipoglycopeptide conjugates to produce alternate vaccines. These vaccines can be used to immunize against diseases caused by H. influenzae when administered to mammals, for example, by the intramuscular or parenteral route, or when delivered using microparticles, capsules, liposomes and targeting molecules, such as toxins or fragments thereof, and antibodies, to cells of the immune system or mucosal surfaces.

(xiv) Utility of D15 as carrier protein for the production of glycoconjugates.

To determine whether D15 may serve both as a protective antigen and a carrier, D15-PRP conjugation experiments were performed as described in Example 14. The D15-PRP conjugates were found to be highly immunogenic in rabbits and able to elicit both anti-D15 and anti-PRP IgG antibody responses as judged by D15-specific ELISA and PRP-BSA immunoassay (Table 9). These results clearly demonstrate the practical utility of D15 as a carrier protein for glycoconjugation technology.

In preferred embodiments of the present invention, the carrier function of D15 can be generally utilized to prepare chimeric molecules and conjugate vaccines against pathogenic bacteria, including encapsulated bacteria. Thus, the glycoconjugates of the present inventions may be applied to vaccinations to confer protection against infection with any bacteria having polysaccharide antigens, including, for example, Haemophilus influenzae, Streptococcus pneumoniae, Escherichia coli, Neisseria meningitidis, Salmonella typhi, Streptococcus mutans, Cryptococcus neoformans, Klebsiella, Staphylococcus aureus and Pseudomonas aeruginosa.

In another embodiment, the carrier function of D15 may be used, for example, to induce immunity toward abnormal polysaccharides of tumor cells, or to produce

anti-tumor antibodies that can be conjugated to chemotherapeutic or bioactive agents.

Accordingly, the present invention provides the primary sequence and the preparation of an antigen (D15) of H. influenzae that can be used in the prevention and diagnosis of diseases caused by Haemophilus. In particular, the inventors discovered that recombinant D15 or its fragments, can elicit protective antibody responses against live H. influenzae type b bacteria challenge. Thus, the present inventions have utility in vaccines. The invention also discloses the nucleotide sequences of the D15 genes isolated from both H. influenzae type b strains and non-typeable isolates. The DNA segments encoding D15 are disclosed and show minor polymorphism in both their nucleotide and derived amino acid sequences (Figures 1F and 3). These DNA segments may be used to provide an immunogen essentially free from other H. influenzae antigens (such as PRP and lipooligosaccharides (LOS)) through the application of recombinant DNA technology. The present disclosure further provides novel techniques which can be employed for preparing essentially pure D15 or fragments thereof, as well as functional analogs. The recombinant D15 protein, fragment or analog thereof, may be produced in a suitable expression system, such as E. coli, Haemophilus, Bordetella, Bacillus, Fungi, Yeast, Baculovirus, Poxvirus, vaccinia or mammalian expression systems.

In one embodiment, the present invention concerns the process of preparing vaccine compositions which include purified recombinant D15 protein (rD15) or rD15 fragments that are immunologically cross-reactive with native D15. In particular, the gene coding the entire D15 protein and a DNA segment encoding an N-terminal rD15 fragment fused to the glutathione-S-transferase gene have been constructed and expressed in E. coli. The expressed

rD15 protein and its fragments were found to cross-react immunologically with the native D15 antigen isolated from both typeable and non-typeable H. influenzae isolates and thus represent cross-reactive immunogens for inclusion in 5 a vaccine against diseases caused by H. influenzae. Furthermore, Haemophilus convalescent serum recognized D15 purified from H. influenzae as described herein, rD15 and N-terminal rD15 fragment.

In another embodiment, the present invention 10 provides a gene coding for the outer membrane protein D15 from H. influenzae having the specific nucleotide sequences described herein or ones substantially homologous thereto (i.e. those which hybridize under stringent conditions to such sequences), for genetically 15 engineering hybrids or chimeric proteins containing a D15 fragment fused to another polypeptide or protein or a polysaccharide, such as H. influenzae outer membrane proteins, for example, P1, P2, or P6 or PRP. As a result, the hybrids, chimeric proteins or glycoconjugates 20 may have higher protectivity against H. influenzae than D15, or P1, or P2, or P6, or PRP alone.

Thus, D15 outer membrane protein can function both as a protective antigen and as a carrier in a conjugate vaccine to provide autologous T-cell priming, wherein the 25 hapten part of the conjugate is the capsular polysaccharide moiety (PRP) of H. influenzae. This D15-carbohydrate conjugate can elicit antibodies against both PRP and D15, and thus should enhance the level of protection against H. influenzae-related diseases, 30 especially in infants.

In another embodiment, the present invention comprises an essentially pure form of at least one protein or peptide containing an amino acid sequence corresponding to at least one antigenic determinant of 35 D15, which peptide is capable of eliciting polyclonal antibodies against H. influenzae in mammals. These D15-

specific antibodies are useful in test kits for detecting the presence of H. influenzae in biological samples. The peptides can have, for example, the amino acid sequences corresponding to residues 20-49, 45-74, 68-99, 93-122, 5 114-143, 135-164, 157-187, 180-209, 199-228, 219-249, 241-270, 262-291, 282-312, 304-333, 325-354, 346-375, 367-396, 390-416, 410-435, 430-455, 450-477, 471-497, 491-516, 511-538, 532-559, 554-582, 577-602, 596-625, 619-646, 641-666, 662-688, 681-709, 705-731, 725-750, 10 745-771, 769-798 (SEQ ID NOS: 14 to 49) of the D15 protein of the H. influenzae type b Ca strain, respectively, as set forth in Table 2 below, or any portion, variant or mutant thereof which retains immunogenicity.

15 In yet another embodiment, the present invention provides pure native D15 protein, extracted and chromatographically purified from cultures of H. influenzae typeable or non-typeable isolates. The novel procedures involves extraction of the D15 protein from 20 cell paste by techniques known for other outer membrane proteins, with an aqueous detergent solution, followed by purification by centrifugation and chromatography. The purified native D15 antigen can be used to immunize mammals against diseases caused by H. influenzae, for 25 example, by the intramuscular or the parenteral routes, or by delivering it using microparticles, capsules, liposomes and targeting molecules, such as toxins or fragments thereof, and antibodies.

Another aspect of the present invention is that the 30 D15 outer membrane protein, fragments or analogs thereof or peptides corresponding to portions of D15 may be components of a multivalent vaccine against otitis media. This multivalent vaccine comprises at least one immunogenic determinant of D15 as described herein, along 35 with at least one protective antigen isolated from Streptococcus pneumoniae, Branhamella (Moroxella)

catarrhalis, Staphylococcus aureus, or respiratory syncytial virus, in the presence or absence of adjuvant.

The D15 peptides (Table 2) or any portion, variant or mutant thereof, can easily be synthesized either 5 manually or with a commercially available peptide synthesizer, such as the Applied Biosystems Model 430A synthesizer.

It is clearly apparent to one skilled in the art, that the various embodiments of the present invention 10 have many applications in the fields of vaccination, diagnosis, and treatment of diseases caused by Haemophilus infections, and the generation of immunological reagents. A further non-limiting discussion of such uses is further presented below.

15 1. Vaccine preparation and use

Immunogenic compositions, suitable for use as vaccines, may be prepared from immunogenic D15 outer membrane protein, fragments or analogs thereof and/or peptides corresponding to portions of D15 as disclosed 20 herein. The vaccine elicits an immune response which produces antibodies, including anti-D15 outer membrane protein antibodies and antibodies against D15 that are opsonizing or bactericidal. Should the vaccinated subject be challenged by Haemophilus, the antibodies bind 25 to the D15 outer membrane protein and thereby inactivate the bacterium. Opsonizing and bactericidal antibodies represent examples of antibodies useful in protection against disease.

Vaccines containing peptides are generally well 30 known in the art, as exemplified by U.S. Patents 4,601,903; 4,599,231; 4,599,230; and 4,596,792; all of which references are incorporated herein by reference. As to any further reference to patents and references in this description, they are as well hereby incorporated by 35 reference without any further notice to that effect. Vaccines may be prepared as injectables, as liquid

solutions or emulsions. The D15 outer membrane protein, fragments or analogs thereof or peptides corresponding to portions of D15 may be mixed with physiologically-acceptable excipients which are compatible with the D15

5 outer membrane protein, fragments, analogs or peptides. Excipients may include, water, saline, dextrose, glycerol, ethanol, and combinations thereof. The vaccine may further contain minor amounts of auxiliary substances such as wetting or emulsifying agents, pH buffering

10 agents, or adjuvants to enhance the effectiveness of the vaccines. Methods of achieving adjuvant effect for the vaccine includes use of agents, such as aluminum hydroxide or phosphate (alum), commonly used as 0.05 to 0.1 percent solution in phosphate buffered saline.

15 Vaccines may be administered parenterally, by injection subcutaneously or intramuscularly. Alternatively, other modes of administration including suppositories and oral formulations may be desirable. For suppositories, binders and carriers may include, for example,

20 polyalkalene glycols or triglycerides. Oral formulations may include normally employed incipients such as, for example, pharmaceutical grades of saccharine, cellulose, magnesium carbonate and the like. These compositions take the form of solutions, suspensions, tablets, pills,

25 capsules, sustained release formulations or powders and contain 10-95% of the D15 outer membrane protein, fragment analogs and/or peptides.

The vaccines are administered in a manner compatible with the dosage formulation, and in an amount which is

30 therapeutically effective, protective and immunogenic. The quantity to be administered depends on the subject to be treated, including, for example, the capacity of the individual's immune system to synthesize antibodies, and if needed, to produce a cell-mediated immune response.

35 Precise amounts of active ingredient required to be administered depend on the judgment of the practitioner.

However, suitable dosage ranges are readily determinable by one skilled in the art and may be of the order of micrograms of the D15 outer membrane protein, analog, fragment and/or peptides. Suitable regimes for initial 5 administration and booster doses are also variable, but may include an initial administration followed by subsequent administrations. The dosage of the vaccine may also depend on the route of administration and varies according to the size of the host.

10 The nucleic acid molecules encoding the D15 outer membrane protein of the present invention may also be used directly for immunization by administration of the DNA directly, for example, by injection for genetic immunization or by constructing a live vector, such as 15 Salmonella, BCG, adenovirus, poxvirus or vaccinia. A discussion of some live vectors that have been used to carry heterologous antigens to the immune system are discussed in, for example, O'Hagan (1992). Processes for the direct injection of DNA into test subjects for 20 genetic immunization are described in, for example, Ulman et al. (1993).

The use of peptides in vivo may first require their chemical modification since the peptides themselves may not have a sufficiently long serum and/or tissue half-life. Such chemically modified peptides are referred to 25 herein as peptide analogs. The term peptide analog extends to any functional chemical equivalent of a peptide characterized by its increased stability and/or efficacy in vivo or in vitro in respect of the practice 30 of the invention. The term peptide analog is also used herein to extend to any amino acid derivative of the peptides as described herein. Peptide analogs contemplated herein are produced by procedures that include, but are not limited to, modifications to side 35 chains, incorporation of unnatural amino acids and/or their derivatives during peptide synthesis and the use of

cross-linkers and other methods which impose conformational constraint on the peptides or their analogs.

Examples of side chain modifications contemplated by 5 the present invention include modification of amino groups, such as by reductive alkylation by reaction with an aldehyde followed by reduction with NaBH<sub>4</sub>; amidation with methylacetimidate; acetylation with acetic anhydride; carbamylation of amino groups with cyanate; 10 trinitrobenzylolation of amino groups with 2, 4, 6, trinitrobenzene sulfonic acid (TNBS); alkylation of amino groups with succinic anhydride and tetrahydrophthalic anhydride; and pyridoxylation of lysine with pyridoxa-5'-phosphate followed by reduction with NaBH<sub>4</sub>.

15 The guanidino group of arginine residues may be modified by the formation of heterocyclic condensation products with reagents such as 2, 3-butanedione, phenylglyoxal and glyoxal.

The carboxyl group may be modified by carbodiimide 20 activation via o-acylisourea formation followed by subsequent derivatisation, for example, to a corresponding amide.

Sulphydryl groups may be modified by methods, such 25 as carboxymethylation with iodoacetic acid or iodoacetamide; performic acid oxidation to cysteic acid; formation of mixed disulphides with other thiol compounds; reaction with maleimide; maleic anhydride or other substituted maleimide; formation of mercurial derivatives using 4-chloromercuribenzoate, 4- 30 chloromercuriphenylsulfonic acid, phenylmercury chloride, 2-chloromercuric-4-nitrophenol and other mercurials; carbamylation with cyanate at alkaline pH.

Tryptophan residues may be modified by, for example, 35 oxidation with N-bromosuccinimide or alkylation of the indole ring with 2-hydroxy-5-nitrobenzyl bromide or sulphonyl halides. Tryosine residues may be altered by

nitration with tetranitromethane to form a 3-nitrotyrosine derivative.

Modification of the imidazole ring of a histidine residue may be accomplished by alkylation with iodoacetic acid derivatives or N-carbethoxylation with diethylpyrocarbonate.

Examples of incorporating unnatural amino acids and derivatives during peptide synthesis include, but are not limited to, use of norleucine, 4-amino butyric acid, 4-amino-3-hydroxy-5-phenylpentanoic acid, 6-aminohexanoic acid, t-butylglycine, norvaline, phenylglycine, ornithine, sarcosine, 4-amino-3-hydroxy-6-methylheptanoic acid, 2-thienyl alanine and/or D-isomers of amino acids.

## 2. Immunoassays

The D15 outer membrane protein, analog, fragment and/or peptides of the present invention are useful as antigens in immunoassays, including enzyme-linked immunosorbent assays (ELISA), RIAs and other non-enzyme linked antibody binding assays or procedures known to the art for the detection of anti-bacterial, Haemophilus, D15 and/or peptide antibodies. In ELISA assays, the D15 outer membrane protein, fragment or analogs thereof and/or peptides corresponding to portions of D15 outer membrane protein are immobilized onto a selected surface, for example, a surface exhibiting a protein affinity, such as the wells of a polystyrene microtiter plate. After washing to remove incompletely adsorbed D15 outer membrane protein, analog, fragment and/or peptides, a nonspecific protein, such as bovine serum albumin (BSA) or casein, that is known to be antigenically neutral with regard to the test sample may be bound to the selected surface. This allows for blocking of nonspecific adsorption sites on the immobilizing surface and thus decreases the background caused by nonspecific bindings of antisera onto the surface. Normally, the peptides

employed herein are in the range of 12 residues and up and preferably 14 to 30 residues.

The immobilizing surface is then contacted with a sample such as clinical or biological materials to be tested in a manner conducive to immune complex (antigen/antibody) formation. This may include diluting the sample with diluents, such as BSA, bovine gamma globulin (BGG) and/or phosphate buffered saline (PBS)/Tween. The sample is then allowed to incubate for from 2 to 4 hours, at temperatures, such as of the order of 25° to 37°C. Following incubation, the sample-contacted surface is washed to remove non-immunocomplexed material. The washing procedure may include washing with a solution such as PBS/Tween, or a borate buffer.

Following formation of specific immunocomplexes between the test sample and the bound D15 outer membrane protein, analog, fragment and/or peptides, and subsequent washing, the occurrence, and even amount, of immunocomplex formation may be determined by subjecting the immunocomplex to a second antibody having specificity for the first antibody. If the test sample is of human origin, the second antibody is an antibody having specificity for human immunoglobulins and, in general, IgG. To provide detecting means, the second antibody may have an associated activity, such as an enzymatic activity that will generate, for example, a color development upon incubating with an appropriate chromogenic substrate. Quantification may then be achieved by measuring the degree of color generation using, for example, a visible spectra spectrophotometer.

### **3. Use of sequences as hybridization probes**

The nucleotide sequences of the present invention, comprising the sequence of the D15 outer membrane protein, now allow for the identification and cloning of the D15 outer membrane protein genes from any species of

Haemophilus and other bacteria that have genes encoding D15 outer membrane proteins.

The nucleotide sequences comprising the sequence encoding the D15 outer membrane protein of the present invention are useful for their ability to selectively form duplex molecules with complementary stretches of other D15 genes. Depending on the application, a variety of hybridization conditions may be employed to achieve varying degrees of selectivity of the probe toward the other D15 genes. For a high degree of selectivity, stringent conditions are used to form the duplexes, such as low salt and/or high temperature conditions, such as provided by 0.02 M to 0.15 M NaCl at temperatures of between about 50°C to 70°C. For some applications, less stringent hybridization conditions are required such as 0.15 M to 0.9 M salt, at temperatures ranging from between about 20°C to 55°C. Hybridization conditions can also be rendered more stringent by the addition of increasing amounts of formamide, to destabilize the hybrid duplex. Thus, particular hybridization conditions can be readily manipulated, and will generally be a method of choice depending on the desired results.

In a clinical diagnostic embodiment, the nucleic acid sequences of the D15 outer membrane protein genes of the present invention may be used in combination with an appropriate means, such as a label, for determining hybridization. A wide variety of appropriate indicator means are known in the art, including radioactive, enzymatic or other ligands, such as avidin/biotin, which are capable of providing a detectable signal. In some diagnostic embodiments, an enzyme tag, such as urease, alkaline phosphatase or peroxidase, instead of a radioactive tag may be used. In the case of enzyme tags, colorimetric indicator substrates are known which can be employed to provide means visible to the human eye or

spectrophotometrically, to identify specific hybridization with samples containing D15 gene sequences.

The nucleic acid sequences of D15 genes of the present invention are useful as hybridization probes in solution hybridizations and in embodiments employing solid-phase procedures. In embodiments involving solid phase procedures, the test DNA (or RNA) from samples, such as clinical samples, including exudates, body fluids (e. g., serum, amniotic fluid, middle ear effusion, sputum, bronchoalveolar lavage fluid) or even tissues, is adsorbed or otherwise affixed to a selected matrix or surface. The fixed, single-stranded nucleic acid is then subjected to specific hybridization with selected probes comprising the nucleic acid sequences of the D15 genes or fragments thereof of the present invention under desired conditions. The selected conditions will depend on the particular circumstances based on the particular criteria required depending on, for example, on the G+C contents, type of target nucleic acid, source of nucleic acid, size of hybridization probe etc. Following washing of the hybridization surface so as to remove non-specifically bound probe molecules, specific hybridization is detected, or even quantified, by means of the label. The selected probe should be at least 18 bp and may be in the range of 30 bp to 90 bp long.

#### 4. Expression of the D15 outer membrane protein genes

Plasmid vectors containing replicon and control sequences which are derived from species compatible with the host cell may be used for the expression of the D15 outer membrane protein genes in expression systems. The vector ordinarily carries a replication site, as well as marking sequences which are capable of providing phenotypic selection in transformed cells. For example, E. coli may be transformed using pBR322 which contains genes for ampicillin and tetracycline resistance and thus provides easy means for identifying transformed cells.

The pBR322 plasmid, or other microbial plasmid or phage must also contain, or be modified to contain, promoters which can be used by the microbial organism for expression of its own proteins.

5 In addition, phage vectors containing replicon and control sequences that are compatible with the host microorganism can be used as a transforming vector in connection with these hosts. For example, the phage in lambda GEM™-11 may be utilized in making recombinant 10 phage vectors which can be used to transform host cells, such as E. coli LE392.

Promoters commonly used in recombinant DNA construction include the  $\beta$ -lactamase (penicillinase) and lactose promoter systems and other microbial promoters, 15 such as the T7 promoter system. Details concerning the nucleotide sequences of promoters are known, enabling a skilled worker to ligate them functionally with plasmid vectors. The particular promoter used generally is a matter of choice depending upon the desired results. 20 Hosts that are appropriate for expression of the transferrin receptor genes, fragment analogs or variants thereof include E. coli, Bacillus, Haemophilus, Bordetella, fungi, yeast, or the baculovirus and poxvirus expression systems may be used.

25 In accordance with an aspect of this invention, it is preferred to make the D15 outer membrane protein, fragment or analog thereof by recombinant methods, particularly since the naturally occurring D15 protein as purified from culture of a species of Haemophilus may 30 include undesired contaminants, including trace amounts of toxic materials. This problem can be avoided by using recombinantly produced D15 outer membrane protein in heterologous systems which can be isolated from the host in a manner to minimize toxins in the purified material. 35 Particularly desirable hosts for expression in this regard include Gram positive bacteria which do not have

lipopolysaccharide (LPS) and are, therefore, endotoxin free. Such hosts include species of Bacillus and may be particularly useful for the production of non-pyrogenic D15 outer membrane protein, fragments or analogs thereof.

5

#### BIOLOGICAL DEPOSITS

Certain plasmids that contain at least a portion coding for a D15 outer membrane protein from strains of Haemophilus influenzae that are described and referred to herein have been deposited with the American Type Culture Collection (ATCC) located at Rockville, Maryland USA pursuant to the Budapest Treaty and prior to the filing of this application. Samples of the deposited plasmids will become available to the public upon grant of a patent based upon this United States patent application. 10  
15 The invention described and claimed herein is not to be limited in scope by plasmids deposited, since the deposited embodiment is intended only as an illustration of the invention. Any equivalent or similar plasmids that encode similar or equivalent antigens as described 20 in this application are within the scope of the invention.

#### DEPOSITE SUMMARY

	Clone	H. influenzae	ATCC Designation	Date Deposited
25	DS-712-2-1	Eagan	75604	November 4, 1993
	JB-1042-5-1	SB33	75606	November 4, 1993
	DS-880-1-2	Eagan	75605	November 4, 1993

The above disclosure generally describes the present 30 invention. A more complete understanding can be obtained by reference to the following specific Examples. These Examples are described solely for purposes of illustration and are not intended to limit the scope of

the invention. Although specific terms have been employed herein, such terms are intended in a descriptive sense and not for purposes of limitations. Immunological and recombinant DNA methods may not be explicitly described 5 in this disclosure but are well within the scope of those skilled in the art.

EXAMPLES

Methods of molecular genetics, protein biochemistry, and immunology used but not explicitly described in this 10 disclosure and these EXAMPLES are amply reported in the scientific literature and are well within the ability of those skilled in the art.

Example 1

This Example illustrates the cloning and sequencing 15 of the D15 genes.

Genomic DNA was purified from the Haemophilus influenzae type b strain Ca by lysis of the bacteria with pronase and sodium dodecylsulphate followed by phenol extraction and isopropanol precipitation, according to 20 Berns and Thomas, 1965. The DNA was then partially digested with EcoRI and the DNA fraction containing 6-10 kb fragments was isolated following electrophoresis in low-melting point agarose. These fragments were ligated into a lambda gt11 Amp1 vector (Thomas and Rossi, 1986) 25 and cloned as a lysogen into E. coli strain BTA282. Recombinant clones were selected for their ampicillin resistance conferred by the vector. To identify clones producing H. influenzae type b antigen, the clones were replica-plated on nitrocellulose filters and duplicate 30 colonies induced for expression by temperature switch to 42°C for 2 hours. Colonies were lysed by wetting the filters with 1% sodium dodecylsulphate (SDS). The filters were then placed into a chloroform-saturated atmosphere for 15 min. The filters were then assayed by colony 35 radioimmuno-assay using a hyperimmune rabbit anti-H. influenzae type b antiserum absorbed with E. coli lysate

for antigen expression. Clones shown by autoradiography to be producing H. influenzae type b antigens were further purified and their replicates retested for reactivity with the hyperimmune anti-H. influenzae type 5 b antiserum. The antiserum absorbed with  $10^{10}$  intact H. influenzae type b bacteria (strain Ca) was used as negative control.

A number of clones were identified which reacted with the unabsorbed, but not with the absorbed antiserum 10 and were further analysed. One of the clones, D15, was purified, grown and found to produce a H. influenzae type b antigen which migrated in sodium dodecyl sulphate polyacrylamide gels with a  $M_r$  of about 80 kDa. Lysates from the D15 clone were coupled to Sepharose™ 4B gel and 15 used to affinity-purify anti-D15 antibodies. This procedure is described by Thomas et al, 1990, except that the apparent  $M_r$  was initially reported to be about 103 kDa. The affinity-purified antibodies to D15 were then shown to react with an  $M_r$  80 kDa protein in an outer 20 membrane protein preparation of H. influenzae type b (sarcosyl insoluble fraction - Carbone et al, 1986). Radioimmuno dot blots and Western blots analyses of membrane preparations from both type b and nontypeable Haemophilus influenzae strains showed that affinity- 25 purified anti-D15 antibodies reacted with all isolates. These antibodies were found to be capable of passively protecting infant rats from bacteraemia following intraperitoneal injection of live H. influenzae type b bacteria. The specificity of the protection was 30 confirmed by absorbing out the protective activity of anti-D15 antibodies with a lysate of E. coli expressing D15 coupled to Sepharose. The protection studies have been described in detail by Thomas et al, 1990.

DNA from the lambda gt11 Amp1 D15 phage was isolated 35 and a 5.7 kb fragment was released by EcoRI digestion. This fragment was subcloned into pUC19 and the resulting

plasmid transformed into E. coli HB101. Recombinant bacteria were found to produce the expected  $M_r$  80 kDa H. influenzae type b antigen when examined by Western blotting. The insert DNA was then characterised by 5 restriction endonuclease mapping. A 2.8 kb HindIII-EcoRI fragment was subcloned into pUC19 to generate plasmid pUC19/D15, which was transformed into E. coli HB101. The recombinant bacteria expressed a  $M_r$  80 kD protein recognized by D15-specific antibodies on Western blot 10 analysis of E. coli lysates.

Plasmid DNA was prepared from two individual colonies of recombinant E. coli HB101 containing the pUC19/D15 plasmid using standard techniques. Oligonucleotide sequencing primers of 17-25 bases in 15 length were synthesized on the ABI model 380B DNA Synthesizer and purified by chromatography using OPC cartridges obtained from Applied Biosystems Inc., and used in accordance with the manufacturers recommendations. Samples were sequenced using the ABI model 370A DNA 20 Sequencer and dye terminator chemistry according to manufacturers' protocols. This sequence analysis indicated that the D15 gene contains an open reading frame encoding for 789 amino acids, including a putative signal sequence (Figure 1). The derived amino acid 25 sequence was found to contain the sequence of an internal peptide obtained by thrombin digestion of native D15 that had been chemically determined. The amino acid composition of D15 derived from the D15 gene sequence was comparable (within experimental error) to that of the 30 native protein as determined by amino acid analysis.

#### Example 2

This Example illustrates the preparation of chromosomal DNA from Haemophilus influenzae strains Eagan, MinnA, SB33, and PAK 12085.

H. influenzae strains were grown on Mueller-Hinton agar or in brain heart infusion broth as described by Harkness et al., 1992.

Eagan chromosomal DNA

5        Bacteria from 50 mL of culture were pelleted by centrifugation at 5,000 rpm, 20 minutes, 4°C. The pellet was resuspended in 25 mL TE (10mM Tris, 1mM EDTA, pH 8.0) and 2 x 5mL aliquots used for chromosomal DNA preparation. To each aliquot were added 0.6 mL of 10%  
10 sarkosyl and 0.15 mL of 20mg/mL proteinase K and the samples incubated at 37°C for 1 hour. The lysate was extracted once with Tris-saturated phenol (pH 8.0) and three times with chloroform:isoamyl alcohol (24:1). The aqueous phase was pooled for a final volume of 7 mL.  
15 Then, 0.7 mL of 3M sodium acetate (pH 5.2) and 4.3 mL of isopropanol were added to precipitate the DNA which was spooled, rinsed with 70% ethanol, dried, and resuspended in 1 mL of water.

MinnA, SB33, and PAK 12085 chromosomal DNA

20       Bacteria from 50 mL of culture were pelleted by centrifugation at 5,000 rpm for 15-20 minutes, at 4°C, in a Sorvall RC-3B centrifuge. The cell pellet was resuspended in 10 mL of TE (10mM Tris-HCl, 1mM EDTA, pH 7.5), pronase was added to 500 µg/mL, and SDS to 1%. The 25 sample was incubated at 37°C for about 4 hours until a clear lysate was obtained. The lysate was extracted once with Tris-saturated phenol, once with Tris-saturated phenol/chloroform (1:1), and once with chloroform. The final aqueous phase was dialysed for 24 hours against 2 x 500 mL of 1M NaCl at 4°C, changing the buffer once, and for 24 hours against 2 x 500 mL of TE at 4°C, changing the buffer once. The final dialysate was aliquotted for subsequent use.

Example 3

35       This Example illustrates the preparation of Haemophilus influenzae chromosomal libraries.

H. influenzae Eagan and PAK 12085 chromosomal DNAs were digested with Sau3A I (0.5 unit/10 µg DNA) at 37°C for 15 minutes and size-fractionated by agarose gel electrophoresis. Gel slices corresponding to DNA fragments of 15-23 kb were excised and DNA was electroeluted overnight in dialysis tubing containing 3 mL of TAE (40mM Tris-acetate, 1mM EDTA, pH 8.0) at 14V. The DNA was precipitated twice and resuspended in water before overnight ligation with EMBL3 BamH I arms (Promega). The ligation mixture was packaged using the Lambda *in vitro* packaging kit (Amersham) according to the manufacturer's instructions and plated onto E. coli NM539 cells. The library was titrated, then amplified and stored at 4°C under 0.3% chloroform.

15 MinnA chromosomal DNA (10 µg) was digested with Sau3A I (40 units) for 2, 4, and 6 minutes then size-fractionated on a 10-30% sucrose gradient in TNE (20mM Tris-HCl, 5mM NaCl, 1mM EDTA, pH 8.0). Fractions containing DNA fragments >5 kb were pooled and precipitated. In a second experiment, chromosomal DNA (2.6 µg) was digested with Sau3A I (4 units) for 1, 2, and 3 minutes and size-fractionated by preparative agarose gel electrophoresis. Gel slices containing DNA fragments of 10-20 kb were excised and DNA extracted by 25 a standard freeze/thaw technique. The size-fractionated DNA from the two experiments was pooled for ligation with BamH I arms of EMBL3 (Promega). The ligation mix was packaged using the Gigapack II packaging kit (Amersham) and plated on E. coli LE392 cells. The library was 30 titrated, then amplified and stored at 4°C under 0.3% chloroform.

SB33 chromosomal DNA (20 µg) was digested with Sau3A I (40 units) for 2, 4, or 6 minutes and size-fractionated on a 10-30% sucrose gradient in TNE (20mM Tris-HCl, 5mM 35 NaCl, 1mM EDTA, pH 8.0). Fractions containing fragments >5 kb were pooled. In a second experiment, SB33

chromosomal DNA (2 µg) was digested with Sau3A I (4 units) for 2, 4, or 6 minutes and size-fractionated on a preparative agarose gel. Gel slices containing DNA fragments of 10-20 kb were excised and DNA extracted by 5 a standard freeze/thaw technique. The size-fractionated DNA from both experiments was pooled for ligation with BamH I arms of EMBL3 (Promega). The ligation mix was packaged using the Gigapack II packaging kit and plated on LE392 cells. The library was titrated, then amplified 10 and stored at 4°C under 0.3% chloroform.

Example 4

This Example illustrates the screening of the DNA libraries.

The Eagan, MinnA, SB33, and PAK 12085 DNA libraries 15 were plated onto LE392 cells on NZCYM plates using 0.7% top agarose in NZCYM as overlay. Plaque lifts onto nitrocellulose filters were performed following standard procedures, and filters were processed and hybridized with a digoxigenin-labelled D15 probe prepared according 20 to the manufacturer's specifications (Boehringer Mannheim). The probe was the EcoR I/Hind III fragment from pUC19/D15 containing the entire Ca D15 gene (Figure 2). Putative plaques were plated and submitted to a second round of screening using the same procedures. 25 Phage DNA was prepared from 500 mL of culture using standard techniques, the insert DNA was excised by Sal I digestion, and cloned into pUC to generate clones DS-712-2-1 (Eagan), DS-691-1-5 (MinnA), JB-1042-5-1 (SB33), and JB-1042-9-4 (PAK 12085), which are shown in Figure 2.

30 The nucleotide sequences of the D15 genes from H. influenzae type b strains Eagan and MinnA the non-typeable H. influenzae strains SB33 and PAK 12085 were determined and compared with that for strain Ca, as seen in figures 1b, 1C, 1D, 1E and 1F. The desired amino acid 35 sequence are shown in Figures 1B, 1C, 1D and 1E and are

compared with the amino acid sequence of the D15 protein of *H. influenzae* type b Ca (Figure 3).

### Example 5

This Example illustrates the expression of rD15  
5 protein in *E. coli*.

A 2.8 kb fragment HindIII-EcoRI was subcloned into pUC19 and this pUC19/D15 plasmid was transformed into E. coli HB101. Upon induction, the positive clones expressed an 80 kDa protein which was recognized by D15-specific antisera on Western blot analysis. A HindIII-Pst I fragment was also subcloned into pUC19 and shown to express a 67 kDa protein. According to the restriction map, this 67 kDa protein corresponded to a C-terminal truncated D15 protein. On Western blot analysis, this truncated D15 was still recognized by the D15-specific antisera.

Plasmids to express the D15 gene of the non-typeable strain SB33 in E. coli were constructed. Plasmid JB-1042-5-1 containing the SB33 D15 gene and its flanking regions, was digested with EcoR I and Hind III and the 3kb D15 insert subcloned into pUC to give plasmid pRY-60-1 (Figure 4). Appropriate oligonucleotides were synthesized to restore the native D15 sequence between the ATG codon of the expression plasmid pT7-7 and the BsrF I site within the D15 gene. These oligonucleotides had the following sequence:

Plasmid pRY-60-1 was digested with EcoR I and BsrF I and the DNA fragment containing most of the D15 gene was purified. pUC was digested with EcoR I and Nde I and the

vector fragment purified. A multi-component ligation between the pUC and D15 fragments and the oligonucleotides generated plasmid DS-860-1-1 which contains a D15 sequence without a promoter. pT7-7 was 5 digested with Nde I and EcoR I and the vector fragment purified. DS-860-1-1 was digested with Nde I and EcoR I and the D15 insert was purified and ligated with the T7-7 vector generating plasmid DS-880-1-2 (Figure 4).

The plasmid constructions were performed using E. 10 coli JM109 as host. For expression, plasmid DS-880-1-2 was transformed into E. coli BL21/DE3, BL21/DE3/pLyss, or JM109/DE3 cells. Transformation of the cells was performed using either calcium chloride-treated competent 15 cells or by electroporation using a BioRad electroporator. Transformed cells were grown in YT, M9, or NZCYM media and induced with IPTG or other inducing agents.

#### Example 6

This Example illustrates the construction and 20 expression of the GST-D15 fragment hybrid gene in E. coli.

A forward sense primer (primer 1) 5'-GGGAATTCCAAAAGATGTTCGT (SEQ ID NO: 52) and a reverse antisense primer CACGAATTCCCTGCAAATC-5' (primer 7 - SEQ 25 ID NO: 53) were used to amplify a 2.8 Kb fragment HindIII-EcoRI of the D15 gene by the polymerase chain reaction that encodes the N-terminal amino acid residues 22 to 223 of the primary sequence of D15 protein (Figure 1A). The nucleotide sequence of the 609bp amplified 30 fragment was confirmed by DNA sequencing. The amplified gene segment was ligated into the pGEX-2T vector downstream from the GST gene and transformed into E. coli TG-1. Colonies expressing the H. influenzae type b antigen were screened with a rabbit anti-H. influenzae 35 type b antiserum by colony radioimmunoassay and isolated. The glutathione-S-transferase-D15 fragment fusion protein

produced by transformed E. coli was isolated by affinity purification on glutathione agarose.

Example 7

This Example describes alternative expression systems for rD15.

The D15 gene or fragments thereof are also expressed in E. coli under the control of other regulated promoters. The D15 gene or fragments thereof are expressed in the absence of the leader peptide, or in other cloning systems where toxicity of D15 expression to the host is not problematic. The gene or fragments thereof are synthesized de novo or by employing the polymerase chain reaction using suitable primers. These genes are cloned into suitable cloning vectors or bacteriophage vectors in E. coli or other suitable hosts directly when toxicity can be avoided. Expression systems are Gram-positive bacteria (such as Bacillus species), pox virus, adenovirus, baculovirus, yeast, fungi, BCG or mammalian expression systems.

20 Example 8

This Example illustrates the protocol for extraction and purification of rD15 from E. coli expression system.

The cell pellet from a 250 mL culture, prepared as described in Example 5, was resuspended in 40 mL of 50 mM Tris, pH 8.0, and disrupted by sonication (3 x 10 min, 70% duty circle). The extract was centrifuged at 20,000 x g and the resulting pellet saved. The initial pellet was re-extracted with 40 mL of 50 mM Tris, 0.5% Triton X-100, 10 mM EDTA, pH 8.0. The suspension was then sonicated for 10 minutes at 70% duty circle. The extract was centrifuged at 300 x g for 5 minutes. The resulting supernatant was centrifuged again at 20,000 x g for 30 min and the resulting pellet was saved. The pellet was resuspended in 50 mM Tris, 0.5% Triton X-100, 10 mM EDTA, pH 8.0. The suspension was then mixed with PBS/ 8 M urea to a final urea concentration of 6 M. The solution was

then dialyzed against PBS to remove urea. After dialysis, the solution was centrifuged at 300 x g for 10 min., the supernatant was saved and stored at 4°C.

Example 9

5        This Example demonstrates the purification of GST-(D15 fragment) fusion protein using glutathione-Sepharose 4B affinity chromatography.

10      Five mg of GST-(D15 fragment) fusion protein crude extract, prepared as described in Example 6, were dissolved in 5 mL of phosphate buffer saline (PBS) containing 1% Triton X-100. The solution was then loaded onto a Glutathione-Sepharose 4B column (2 mL) equilibrated with PBS containing 1% Triton X-100. The run-through of the column was discarded. The column was 15 washed with 20 mL of PBS and the GST-(D15 fragment) fusion protein was eluted with 50 mM Tris-HCl buffer, pH 8.0, containing 5 mM glutathione. Elution was monitored by absorbance at 280 nm. Protein-containing fractions (2 mL/fraction) were collected and pooled. The purity of the 20 protein was assessed by SDS-PAGE (Figure 9, lane 3). The final volume of the purified fusion protein was 6 mL.

Example 10

25      This Example illustrates the protocol used for thrombin digestion of proteins to release the truncated D15 molecule.

The GST-(D15 fragment) fusion protein sample from Example 9 (0.1 to 0.5 mg protein/mL) was dialyzed against 1 L of 50 mM Tris-HCl buffer (pH 8.5) 3 times with at least 2 hour intervals at 4°C to remove protease 30 inhibitors. After dialysis, the solution was treated with human thrombin (Sigma) at a ratio of 1 mL of solution to 25 units of thrombin. The cleavage reaction was carried out at 37°C for 2 hr and analysed by SDS-PAGE (Figure 9, lane 4). The reaction was stopped by placing 35 the solution in ice.

Example 11

This Example illustrates the procedure used for N-terminal rD15 fragment purification from GST using Glutathione-Sepharose 4B affinity chromatography.

A thrombin-digested GST-(D15 fragment) sample, 5 prepared as described in Example 10, was loaded onto a Glutathione-Sepharose 4B column (2 mL) equilibrated with PBS containing 1% Triton X-100. The run-through of the column containing the N-terminal rD15 fragment was saved. After washing the column with 20 mL of PBS, the affinity 10 column was regenerated by removing GST using 50 mM Tris-HCl buffer, pH 8.0, containing 5 mM glutathione. The purity of rD15 fragment was analysed by SDS-PAGE (Figure 9, lane 5). This N-terminal rD15 fragment contains amino acids 63-223 of the D15 protein as a result of cleavage 15 at the spacious thrombin site shown in Figure 1A.

Example 12

This Example illustrates the protocol used for the purification of D15-specific polyclonal antibodies by affinity chromatography using GST-(D15 fragment) fusion 20 protein.

The recombinant GST-(D15 fragment) fusion protein, prepared as described in Example 9, was conjugated to cyanogen bromide-activated Sepharose. The affinity column was then used to purify antibodies from a rabbit 25 hyperimmune anti-*H. influenzae* type b antiserum. The affinity purified-antibodies were shown by immunoblotting to react with a 80 kDa component present in the lysates of *E. coli* transformed with pUC9/D15 and in the lysates of several typeable and nontypeable *H. influenzae* 30 isolates. These results confirmed that the DNA segment encoding the D15 fragment of the fusion protein was part of the open reading frame of the D15 gene.

Similarly, antisera raised against the recombinant fusion protein (Example 9) or the purified N-terminal 35 rD15 fragment (Example 11) reacted with the D15 protein produced by *H. influenzae* strains (Example 13).

Example 13

This Example describes the protocol used for the purification of native D15 from H. influenzae.

Cell paste of the non-typeable H. influenzae SB33  
5 strain, prepared from a culture grown in brain heart infusion medium supplemented with NAD (2 $\mu$ g/mL) and HEMIN (2 $\mu$ g/mL) at 37°C, as described in Panezutti, et al, 1993, was resuspended in 50 mM Tris-HCl, pH 8.0, containing 0.5% Triton X-100 and 10 mM EDTA (20 mL per 1 g of cell  
10 paste). The mixture was stirred at room temperature for 2 hr, then centrifuged at 20,000 x g for 30 minutes. The D15 was located in the supernatant and further purified.

Purification of native D15 was achieved by affinity chromatography using a D15-specific monoclonal antibody  
15 (see Example 24). The D15 extract (25 mL) was mixed with the affinity matrix (1 mL) at room temperature for 2 hr. The mixture was packed into a column and the run-through fraction was discarded. The column was washed sequentially with the following buffers: 50 mM Tris-HCl, pH 8.0, containing 0.5% Triton X-100 and 10 mM EDTA; 1 M HEPES buffer, pH 6.8; 50 mM Tris-HCl, pH 8.0, containing 0.5% Triton X-100 and 10 mM EDTA; and 10 mM phosphate buffer, pH 8.0. D15 was then eluted from the column with 3 mL of 50 mM diethylamine, pH 12.0 and the protein  
20 solution was neutralized by 1 M HEPES, pH 6.8 (1/10 volume). The affinity-purified native D15 was analysed by SDS-PAGE and stored at -20°C.  
25

Example 14

This Example describes the procedure used for the preparation of D15-PRP conjugates.  
30

Haemophilus influenzae type b oligosaccharides (PRP) prepared by controlled acid hydrolysis were conjugated either with the purified native (Example 13) or recombinant D15 (Example 8) as well as with its fragments  
35 (Example 11) using periodate oxidation as described in US Patent 4356170 and further details of which are presented

in Example 17. The mean molecular size of the PRP molecules used for conjugation was determined as being approximately 20,000 Daltons. The conjugation was carried out without a linker molecule but may also be 5 carried out with a linker molecule. A PRP/D15 molar ratio of approximately 7 was used to provide an excess of PRP hapten.

The PRP/rD15 conjugate was tested according to the protocol of Example 18 for immunogenicity in rabbits and 10 elicited both primary and secondary anti-PRP IgG and anti-D15 antibody responses (Table 9). Rabbit anti-rD15-PRP antisera also strongly reacted with both native D15 and rD15 as judged by immunoblot analysis. These data indicate that rD15 can be used as a carrier protein in a 15 conjugate vaccine. In addition, a rD15-PRP conjugate vaccine should ensure a more consistent protection against H. influenzae type b disease, particularly in infants, as a result of the additional homotypic protection provided by antibodies directed against the 20 D15 protein.

#### Example 15

This Example describes the preparation of D15 peptides.

D15 peptides (Table 2) were synthesized using an ABI 25 430A peptide synthesizer and optimized t-Boc chemistry as described by the manufacturer, then cleaved from the resin by hydrofluoric acid (HF). The peptides were purified by reversed-phase high performance liquid chromatography (RP-HPLC) on a Vydac C4 semi-preparative 30 column (1 x 30 cm) using a 15 to 55% acetonitrile gradient in 0.1% trifluoromethyl acetic acid (TFA) developed over 40 minutes at a flow rate of 2 mL/min. All synthetic peptides (Table 2) used in biochemical and immunological studies were >95% pure as judged by 35 analytical HPLC. Amino acid composition analyses of

these peptides performed on a Waters Pico-Tag system were in good agreement with their theoretical compositions.

Example 16

This Example describes the protocol used for D15 peptide-specific antisera production.

Guinea pigs and rabbits were immunized with individual peptides (50 to 200 µg) emulsified with Freund's complete adjuvant and injected intramuscularly. After two booster doses with the same amount of peptide in incomplete Freund's adjuvant at +14 and +28 days, the anti-peptide antisera were collected on day +42 and tested by ELISAs and immunoblotting. Both rabbit and guinea pig antisera were shown to be monospecific for their respective immunizing peptides by the peptide-specific ELISAs (Table 6). In addition, both guinea pig and rabbit antisera raised against D15 peptides reacted with both *H. influenzae* type b and non-typeable D15 on immunoblot analyses. Since most D15 peptides induced strong anti-peptide antibody responses in at least one animal species, they are appropriate immunogens to be included in immunogenic compositions including vaccine preparations.

Example 17

This Example describes the procedure used for the preparation of PRP-BSA conjugates.

0.5 mL of periodate-oxidized PRP (25mg in 1 mL of 0.1 M sodium phosphate buffer, pH 6.0), prepared from native PRP treated with aqueous periodic acid (Carlone et al, 1986), was added to bovine serum albumin (BSA) (1.32 mg ; 0.02 µmol) in 0.5 mL of 0.2 M sodium phosphate buffer, pH 8.0, followed by the addition of sodium cyanoborohydride (14 µg ; 0.22 µmol ; 10 eqv. to BSA). After incubation at 37°C for 5 days, the reaction mixture was dialysed against 4 L of 0.1 M phosphate buffer, pH 7.5. The resulting solution was applied onto an analytical Superose 12 column (15 x 300 mm, Pharmacia)

equilibrated with 0.2 M sodium phosphate buffer, pH 7.2, and eluted with the same buffer. Fractions were monitored for absorbance at 230 nm. The first major protein peak was pooled and concentrated in a Centriprep 30 to 2.2 mL.

5 The amount of protein was determined using the Bio Rad protein assay, and was found to be 300 µg/mL. The presence of PRP in the protein conjugate fraction was confirmed by the Orcinol test.

Example 18

10 This Example describes the protocol used for the production of anti-PRP antisera in animals using rD15-PRP conjugates.

Rabbits were immunized intramuscularly with rD15-PRP conjugates (Example 14) (5 to 50 µg PRP equivalent) mixed  
15 with 3 mg AlPO<sub>4</sub> per mL, followed by two booster doses (half amount of the same immunogen) at 2 week intervals. Antisera were collected every 2 weeks after the first injection, heat-inactivated at 56°C for 30 minutes and stored at -20°C.

20 Example 19

This Example illustrates the reactivity between D15 peptides and anti-peptide and D15-specific antisera using D15-specific and peptide-specific ELISAs.

Microtiter wells (Nunc-Immunoplate, Nunc, Denmark)  
25 were coated with 200 ng of purified rD15 or 500 ng of individual peptides in 50 µL of coating buffer (15 mM Na<sub>2</sub>CO<sub>3</sub>, 35 mM NaHCO<sub>3</sub>, pH 9.6) for 16 hours at room temperature. The plates were then blocked with 0.1% (w/v) BSA in phosphate buffer saline (PBS) for 30 minutes at  
30 room temperature. Serially diluted antisera were added to the wells and incubated for 1 hour at room temperature. After removal of the antisera, the plates were washed five times with PBS containing 0.1% (w/v) Tween-20 and 0.1% (w/v) BSA. F(ab')<sub>2</sub> fragments from goat anti-rabbit,  
35 guinea pig, mouse, or human IgG antibodies conjugated to horseradish peroxidase (Jackson ImmunoResearch Labs Inc.,

PA) were diluted (1/8,000) with washing buffer, and added onto the microtiter plates. After 1 hr incubation at room temperature, the plates were washed five times with the washing buffer. The plates were then developed using 5 the substrate tetramethylbenzidine (TMB) in H<sub>2</sub>O<sub>2</sub> (ADI, Toronto). The reaction was stopped with 1N H<sub>2</sub>SO<sub>4</sub> and the optical density was measured at 450 nm using a Titretek Multiskan II (Flow Labs., Virginia). Two irrelevant peptides as negative controls in the peptide-specific 10 ELISAs. Assays were performed in triplicate, and the reactive titer of each antiserum was defined as the dilution consistently showing 2-fold increase absorbance value over those obtained from the negative controls. The results obtained are summarized in Tables 3, 6 and 8 and 15 in the DETAILED DESCRIPTION OF THE INVENTION above.

Example 20

This Example illustrates the measurement of the anti-PRP IgG titers in rabbit anti-PRP-D15 conjugate antisera using a PRP-specific ELISA.

20 Microtiter wells (Nunc-Immunoplate, Nunc, Denmark) were coated with 200 ng of purified PRP-BSA (see Example 17) in 200 µL of coating buffer (15 mM Na<sub>2</sub>CO<sub>3</sub>, 35 mM NaHCO<sub>3</sub>, pH 9.6) for 16 hours at room temperature. The plates were then blocked with 0.1% (w/v) BSA in phosphate 25 buffer saline (PBS) for 30 minutes at room temperature. Serially diluted rabbit antisera raised against PRP-D15 conjugates were added to the wells and incubated for 1 hour at room temperature. After removal of the antisera, the plates were washed five times with PBS containing 30 0.1% (w/v) Tween-20 and 0.1% (w/v) BSA. F(ab')<sub>2</sub> fragment from goat anti-rabbit IgG antibodies conjugated to horseradish peroxidase (Jackson ImmunoResearch Labs Inc., PA) were diluted (1/8,000) with washing buffer, and added onto the microtiter plates. After 1 hour incubation at 35 room temperature, the plates were washed five times with the washing buffer. The plates were then developed using

the substrate tetramethylbenzidine (TMB) in H<sub>2</sub>O<sub>2</sub> (ADI, Toronto). The reaction was stopped with 1N H<sub>2</sub>SO<sub>4</sub> and the optical density measured at 450 nm using a Titretek Multiskan II (Flow Labs., Virginia). A standard anti-PRP 5 antiserum of known titer was included as positive control. Assays were performed in triplicate, and the reactive titer of each antiserum was defined as the reciprocal of the dilution consistently showing a 2-fold increase in O.D. value over that obtained with the pre- 10 immune serum (Table 9).

Example 21

This Example describes the protocol used for the production of D15-specific antisera using purified D15, rD15 or N-terminal rD15 fragment.

15 New Zealand White rabbits (Maple Lane) and guinea pigs (Charles River) were immunized intramuscularly (IM) with a 10 µg dose of either affinity-purified native D15 (Example 13), recombinant D15 (Example 8) or N-terminal rD15 fragment (Example 11) emulsified in Freund's 20 complete adjuvant (Difco). Animals were boosted on day 28 with another 10 µg dose of affinity-purified D15 or rD15 or rD15 fragment emulsified in Freund's incomplete adjuvant and bled on day 42 via the marginal ear vein. D15-specific polyclonal antibodies were purified from 25 this material as described in Example 12.

Example 22

This Example illustrates the protective activity of D15-specific antisera against H. influenzae type b challenge using the infant rat model of bacteremia.

30 Five-day old infant rats were inoculated subcutaneously (SC) on the dorsum with 0.15 mL of two different rabbit anti-N-terminal rD15 fragments. Pre-immune sera were used as negative controls. One day after immunization, the infant rats were injected 35 intraperitoneally (IP) with 200 colony-forming units (cfu) of Haemophilus influenzae type b Minn A strain (0.1

ml) freshly grown in brain heart infusion (BHI) medium supplemented with cofactors and diluted in PBS containing 0.5 mM MgCl<sub>2</sub> and 0.15 mM CaCl<sub>2</sub>. One day later, blood samples were collected via cardiac puncture under 5 methoxyflurane anaesthesia and plated on chocolate agar plates. The number of bacteria per mL of blood was quantified after 24 hr. The statistical significance of differences observed in the levels of bacteremia relative to controls was analyzed by the Student's t-test. The 10 results are summarized in Table 1.

Example 23

This Example describes the protocol used for the generation of D15-specific T-cell lines.

BALB/c (H-2<sup>d</sup>) mice purchased from Charles River 15 Animal Farm (Montreal, Canada) were individually primed subcutaneously with 20 µg of rD15 adsorbed to 1.5 mg of aluminium phosphate (alum). The animals were boosted twice with the same dose of immunogen at 3 week intervals. Ten days after the final boost, spleens of 20 immunized mice were removed. Splenocytes were cultured at 5.75 x 10<sup>5</sup> cells per well in a final volume of 200 µL of RPMI 1640 medium (Flow Lab.) supplemented with 10% heat-inactivated fetal calf serum (Gibco), 2 mM L-glutamine (Flow Lab.), 100 U/mL penicillin (Flow Lab.) and 5 x 10<sup>-5</sup> 25 M 2-mercaptoethanol (Sigma) in the presence of varying concentrations (1, 10 and 100 µg per mL) of individual D15 peptides (Table 2) in 96-well plates (Nunc, Denmark). Cultures were kept in a humidified incubator in the presence of 5% CO<sub>2</sub>/air. Triplicate cultures were 30 performed for each concentration of each peptide. Five days later, 150 µL of 10% rat concanavalin A culture supernatant diluted in culture medium was added to the microtiter plate wells as a source of Interleukin-2 (IL-2) to expand peptide-specific T-cells. Six days later, 35 150 µL of supernatant were removed from each microculture, and 150 µL of fresh IL-2 containing culture

supernatant added to further expand and maintain the viability of the peptide-specific T-cells. After a further 6 day-incubation, the cells were washed three times, each time with 200 µL of culture medium.

5        Each set of cultures was then stimulated with the corresponding concentrations (1, 10 and 100 µg per mL) of the peptide in the presence of  $2 \times 10^5$  irradiated (1,500 rad) BALB/c spleen cells in a final volume of 200 µL of culture medium. Sixty µL of supernatant were then removed  
10      from each microculture. The supernatants from each triplicate cultures set were pooled. All supernatants were assayed for IL-2, Interleukin-4 and Interferon-gamma (IFN- $\gamma$ ). Detections of IL-2 and IL-4 were performed using murine IL-2 and IL-4 ELISA kits purchased from  
15      Endogen Inc. (MA, USA) respectively. Assay of IFN- $\gamma$  was performed using a mouse IFN- $\gamma$  ELISA kit supplied by Genzyme Corporation (MA, USA). Test culture supernatants were assayed at 1 in 5 dilution according to the manufacturers' instructions. The results obtained are  
20      set forth in Table 7.

Example 25

This Example describes the general procedure used for the production of murine D15-specific monoclonal antibodies.

25        BALB/c mice were immunized intraperitoneally with 20 to 50 µg of the N-terminal rD15 fragment (Example 11) emulsified in Freund's complete adjuvant. Two weeks later, the mice were given another injection of the same amount of immunogen in incomplete Freund's adjuvant  
30      (IFA). Three days before the fusion, the mice were boosted again with the same amount of immunogen in IFA. Hybridomas were produced by fusion of splenic lymphocytes from immunized mice with non-secreting Sp2/0 myeloma cells as previously described by Hamel et al. (1987).  
35      D15-specific hybridomas were cloned by sequential limiting dilutions and screened for anti-D15 monoclonal

antibody production. Eight D15-specific hybridoma cell lines were identified, expanded and frozen in liquid nitrogen. One of the hybridoma cell lines, 6C8-F6-C6, has been partially characterized. The monoclonal antibody 5 (MAb 6C8-F6-C6) reacts with peptide D15-P8. This MAb 6C8-F6-C6 was used to prepare the D15-specific MAb affinity column and purify native D15 from H. influenzae cell paste (Example 13).

TABLE 1

PROTECTIVE EFFECT OF PASSIVELY TRANSFERRED ANTI-N-TERMINAL RD15  
FRAGMENT ANTIBODIES IN THE INFANT RAT MODEL OF BACTEREMIA<sup>1</sup>

Rabbit antisera	cfu/0.1 mL blood		
	Pre-immune	Post-immunization	p value
Rb#434	510 (6/6) <sup>2</sup>	6 (1/6)	<0.001
Rb#435	910 (4/4)	6 (1/4)	<0.001

<sup>1</sup> Five-day old infant rats were passively immunized with 0.15 mL of rabbit anti-N-terminal rD15 fragment s.c. One day later, the infant rats were challenged with 200 cfu of H. influenzae type b strain MinnA (0.1 mL, IP). The blood samples were taken from each rat 24 hours after the challenge and analysed for bacteria counts.

<sup>2</sup> The parentheses indicate the number of rats found to be bacteremic out of the total number of rats challenged.

TABLE 2

SEQUENCE OF OVERLAPPING SYNTHETIC PEPTIDES ENCOMPASSING  
THE ENTIRE D15 ANTIGEN SEQUENCE

PEPTIDES	RESIDUES	SEQUENCES	SEQ ID NO:
D15-P1	20-49	APFVAKDIRVDGVQGDLEQQIRASLPVRAG	14
D15-P2	45-74	PVRAGQRVTDNDVAMIVRSLFVSGRFDDVK	15
D15-P3	68-99	GRFDDVKAHQEGDVLVVSVVAKSIIISDVKIKG	16
D15-P4	93-122	SDVKIKGNNSVIPTEALKQNLDANGFKVGDV	17
D15-P5	114-143	ANGFKVGDVLIKEKLNEFAKSVKEHYASVG	18
D15-P6	135-164	VKEHYASVGRYNATVEPIVNTLPNNRAEIL	19
D15-P7	157-187	PNNRAEILIQINEDDKAKLASLTFKGNESVS	20
D15-P8	180-209	FKGNESVSSSTLQEQMELQPDSWWKKLWGNK	21
D15-P9	199-228	PDSWWKLWGNKFEGAQFEKDLQSIRDYYLN	22
D15-P10	219-249	LQSIRDYYLNNGYAKAQITKTDVQLNDEKT	23
D15-P11	241-270	VQLNDEKTKVNVTIDVNEGLQYDLRSARI	24
D15-P12	262-291	YDLRSARIIGNLGGMSAELPLLSALHLND	25
D15-P13	282-312	PLLSALHLNDTFRRSDIADVENAIKAKLGER	26
D15-P14	304-333	AIKAKLGERGYGSATVNSVPDFDDANKTLA	27
D15-P15	325-354	FDDANKTLAITLVVDAGRRLTVRQLRFEGN	28
D15-P16	346-375	VRQLRFEGNTVSADSTLRQEMRQQEGTWYN	29
D15-P17	367-396	RQQEGTWYNSQLVELGKIRLDRTGFFETVE	30
D15-P18	390-416	GFFETVENRIDPINGSNDEVDVVYKV	31
D15-P19	410-435	DVYYKVKERNNTGSINFFIGYGTESGI	32
D15-P20	430-455	GTESGISYQASVKQDNFLGTGAAVSI	33
D15-P21	450-477	GAAVSIAGTKNDYGT SVN LGY TE PYFTK	34
D15-P22	471-497	TEPYFTKDGVS LGGNV FFEN YDNSKSD	35
D15-P23	491-516	YDNSKSDTSSNYKRTTYGSNVTLGFP	36
D15-P24	511-538	VTLGFPVNENNNSYYVGLGHTYNKISNF	37
D15-P25	532-559	YNKISNF ALEYNRNLYIQSMKFKGNGIK	38
D15-P26	554-582	KGNGIKTNDFDFSFGWNYN SLN RGY FPTK	39
D15-P27	577-602	GYFPTKGVKASLGGRVTIPGSDNKYYK	40
D15-P28	596-625	SDNKYYKLSADVQGFYPLDRDHLWVVS A K	41

D15-P29	619-646	LWVVSAKASAGYANGFGNKRLPFYQTYT	42
D15-P30	641-666	FYQTYTAGGIGSLRGFAYGSIGPNAI	43
D15-P31	662-688	GPNAIYAEYGNNGSGTGTFKKISSDVIG	44
D15-P32	681-709	KISSDVIGGNAIATASAELIVPTPFVSDK	45
D15-P33	705-731	FVSDKSQNTVRTSLFVDAASVWNTKWK	46
D15-P34	725-750	VWNTKWKSDKNGLESDVLKRLPDYGK	47
D15-P35	745-771	LPDYGKSSRIRASTGVGFQWQSPIGPL	48
D15-P36	769-798	GPLVFSYAKPIKKYENDDVEQFQFSIGGSF	49

TABLE 3

REACTIVITY OF RABBIT AND GUINEA PIG ANTI-N-TERMINAL rD15  
FRAGMENT ANTISERA WITH D15 SYNTHETIC PEPTIDES

Peptides	Reactive Titers					
	Rabbit antisera		Guinea pig antisera			860
	3434	435	858	859		
D15-P1	400	1,600	6,400	6,400	6,400	
D15-P2	1,600	<100	100	100	<100	
D15-P3	400	<100	100	<100	<100	
D15-P4	25,600	6,400	<100	<100	<100	
D15-P5	6,400	400	1,600	25,600	400	
D15-P6	1,600	6,400	400	6,400	6,400	
D15-P7	6,400	1,600	25,600	25,600	25,600	
D15-P8	6,400	6,400	25,600	409,600	409,600	
D15-P9	<100	<100	400	1,600	1,600	
D15-P10	<100	<100	400	6,400	<100	

TABLE 4

## INHIBITION OF ANTI-N-TERMINAL rD15 FRAGMENT ANTIBODY-INDUCED PROTECTION BY D15 PEPTIDES IN THE INFANT RAT MODEL OF BACTEREMIA

Group #	Antibody	cfu / 10 $\mu$ L blood	cfu in each group/ cfu in group #4 (control) (%)
1	Anti-D15 Ab + PBS	60 ± 120 (3/7)	3
2	Anti-D15 Ab + peptides	300 ± 240 (6/7)	13
3	Anti-D15 Ab + rD15	1,520 ± 1,280 (7/7)	64
4	PBS + peptides	2,360 ± 1,200 (6/7)	100

One half mL of rabbit anti-N-terminal rD15 fragment antiserum (Anti-rD15 fragment Ab) was mixed with either nine D15 peptides (100  $\mu$ g of peptides D15-P2 to D15-P10, See TABLE 2) or with 600  $\mu$ g of N-terminal rD15 fragment at room temperature for 1 hr. Antiserum and peptides mixed with PBS were used as controls. Seven-day old infant rats were injected s.c. with 0.2 mL of the various preparations. After 24 h, the infant rats were challenged I.P. with 200 cfu of *H. influenzae* type b strain MinnA. The blood samples were taken at 24 h after the challenge. The numbers in parentheses indicate the number of animals that were bacteremic out of the total number of animals challenged. The level of bacteremia is expressed as the mean of values obtained from seven infant rats tested individually ± one standard deviation (SD).

TABLE 5.

INHIBITION OF THE IMMUNOPROTECTION ABILITY OF THE RABBIT  
ANTI-N-TERMINAL rD15 FRAGMENT ANTISERUM ABSORBED WITH D15 PEPTIDES  
(D15-P4 TO D15-P8) IN THE INFANT RAT MODEL OF BACTEREMIA

Group #	Antibody	cfu / 10 $\mu$ L blood	cfu in each group/ cfu in group #3 (%)
1	rD15 Ab + PBS	220 $\pm$ 360 (3/6)	8
2	rD15 Ab + peptides	2,960 $\pm$ 560 (6/6)	106
3	PBS + peptides	2,800 $\pm$ 360 (6/6)	100

One half mL of rabbit anti-rD15 fragment antiserum (rD15 Ab) was mixed with five D15 peptides (peptides P4 to P8, 250  $\mu$ g of each peptide) at room temperature for 1 hr. Antiserum and peptides diluted in PBS were used as controls. Seven-day old infant rats were injected s.c. with 0.2 mL of the indicated material. After 24 h, the infant rats were challenged I.P. with 200 cfu of *H. influenzae* type b strain MinnA. The blood samples were collected 24 h after challenge. The numbers in parentheses indicate the number of animals that were bacteremia out of the total number of animals challenged. The level of bacteremia is expressed as the mean of values obtained from six infant rats tested individually  $\pm$  one SD.

TABLE 6

REACTIVITY OF RABBIT, GUINEA PIG AND MOUSE ANTI-rD15 ANTISERA  
WITH D15 PEPTIDES

Peptide	Rabbit <sup>2</sup>	Reactive Titer <sup>1</sup> Guinea Pig <sup>3</sup>	Mouse <sup>4</sup>
D15-P1	-	-	+
D15-P2	-	+++	+
D15-P3	-	-	+
D15-P4	+	+	+
D15-P5	-	-	+
D15-P6	-	+	+
D15-P7	-	-	+
D15-P8	-	++++	++++
D15-P9	-	-	+
D15-P10	-	-	+++
D15-P11	-	-	+++
D15-P12	-	-	+
D15-P13	-	-	+
D15-P14	+++	+	+
D15-P15	-	-	+
D15-P16	-	-	+
D15-P17	-	-	+
D15-P18	-	-	+
D15-P19	-	-	+
D15-P20	-	-	+
D15-P21	-	-	+
D15-P22	-	-	+
D15-P23	-	-	+
D15-P24	-	-	+
D15-P25	-	-	+
D15-P26	-	-	+++
D15-P27	-	+	+

TABLE 6 (continued)

D15-P28	-	-	+
D15-P29	-	-	+
D15-P30	-	-	+
D15-P31	-	-	+
D15-P32	-	-	+
D15-P33	-	-	+
D15-P34	-	-	+
D15-P35	-	-	+
D15-P36	++++	-	+

<sup>1</sup> The reactive titer is based on peptide-specific ELISAs. +, ++, +++, and ++++ represent reactive titers of animal antisera tested at 1/300, 1/1000, 1/2000, and 1/5000 dilutions, respectively; - means nonreactive.

<sup>2</sup> Titer represents the average value of two rabbit antisera raised against rD15.

<sup>3</sup> Titer represents the average value of two guinea pig antisera raised against rD15.

<sup>4</sup> Titer represents the average value of five mouse antisera raised against r15.

TABLE 7  
T-CELL STIMULATORY ACTIVITY OF D15 PEPTIDES

Peptide	IL-2 <sup>2</sup>	CYTOKINE RELEASE (pg/mL) <sup>1</sup> $\gamma$ -IFN <sup>3</sup>	IL-4 <sup>4</sup>
D15-P1	-	-	-
D15-P2	122	-	-
D15-P3	25	-	-
D15-P4	-	-	-
D15-P5	742	38,000	13
D15-P6	-	-	-
D15-P7	-	-	-
D15-P8	-	-	-
D15-P9	-	-	-
D15-P10	108	1,900	-
D15-P11	-	-	-
D15-P12	1,052	6,100	-
D15-P13	105	6,200	56
D15-P14	-	-	-
D15-P15	-	-	-
D15-P16	48	-	-
D15-P17	-	-	-
D15-P18	32	4,800	-
D15-P19	882	24,500	-
D15-P20	-	-	-
D15-P21	-	-	-
D15-P22	-	-	-
D15-P23	78	-	-
D15-P24	103	-	-
D15-P25	-	-	-
D15-P26	572	6,700	-
D15-P27	274	7,505	68

TABLE 7 (continued)

D15-P28	<b>142</b>	<b>742</b>	-
D15-P29	-	-	-
D15-P30	-	-	-
D15-P31	-	-	-
D15-P32	-	-	-
D15-P33	-	-	-
D15-P34	<b>82</b>	<b>603</b>	-
D15-P35	<b>107</b>	<b>751</b>	-
D15-P36	-	-	-

<sup>1</sup> Results are expressed as mean values of triplicate cultures. All standard deviations were less than 15%. Immunodominant Th1-cell epitopes are highlighted with bold and Th0-cell epitopes are in italics.

TABLE 8

## RABBIT AND GUINEA PIG ANTIBODY RESPONSES TO D15 PEPTIDES

Immunogen	Peptide-specific ELISAs	
	Rabbit <sup>2</sup>	Reactive Titer <sup>1</sup>
D15-P1	102,400	819,200
D15-P2	204,800	1,637,400
D15-P3	51,200	1,637,400
D15-P4	204,800	819,200
D15-P5	51,200	1,637,400
D15-P6	51,200	409,600
D15-P7	204,800	819,200
D15-P8	51,200	409,600
D15-P9	102,400	409,600
D15-P10	102,400	819,200
D15-P11	51,200	819,200
D15-P12	102,400	204,800
D15-P13	NT <sup>4</sup>	204,800
D15-P14	NT	409,600
D15-P15	NT	204,800
D15-P16	NT	819,200
D15-P17	NT	204,800
D15-P18	NT	312,500
D15-P19	NT	312,500
D15-P20	NT	62,500
D15-P21	NT	62,500
D15-P22	NT	12,500
D15-P23	NT	1,562,500
D15-P24	NT	312,500
D15-P25	NT	62,500

TABLE 8 (continued)

D15-P26	NT	500
D15-P27	NT	1,500
D15-P28	NT	1,250
D15-P29	NT	<500
D15-P30	NT	<500
D15-P31	NT	<500
D15-P32	NT	12,500
D15-P33	NT	12,500
D15-P34	NT	62,500
D15-P35	NT	1,250
D15-P36	NT	12,500

<sup>1</sup> The reactive titer is based on peptide-specific ELISAs. A titer below 500 indicates that the peptide is not immunogenic.

<sup>2</sup> Titters represent the average value of obtained for two rabbit antisera raised against the D15 peptide.

<sup>3</sup> Titters represent the average value obtained for two guinea pig antisera raised against the D15 peptide.

<sup>4</sup> NT: not tested.

TABLE 9  
RABBIT IgG ANTIBODY RESPONSE TO D15-PRP CONJUGATE

Rabbit <sup>1</sup> #	Reactive Titer Against <sup>2</sup>			
	PRP		rD15	
	2 doses	3 doses	2 doses	3 doses
489-1	1,600	3,200	1,600	6,400
490-1	1,600	1,600	6,400	25,600

<sup>1</sup> Rabbits were immunized intramuscularly with rD15-PRP conjugates (5 to 50 µg PRP equivalent) mixed with 3 mg AlPO<sub>4</sub> per mL, followed by two booster doses (half amount of the same immunogen) at 2 week intervals.

<sup>2</sup> Reactive titres is based on PRP specific and D-15 specific ELISAs.

SUMMARY OF THE DISCLOSURE

In summary of this disclosure, the present invention provides purified and isolated nucleic acid molecules containing genes encoding the D15 outer membrane protein, 5 the sequences of these genes and the derived amino acid sequences thereof. The invention also provides peptides corresponding to portions of the D35 outer membrane protein. In addition, the invention provides antibodies raised against D15 outer membrane protein, fragments and 10 peptides. The genes, DNA sequences, antibodies and peptides are useful for diagnosis, immunization and the generation of diagnostic and immunological reagents. Vaccines based on expressed recombinant D35, portions thereof or peptides derived from the provided sequences 15 can be prepared for prevention of H. influenzae disease. Modification are possible within the scope of the invention.

REFERENCES

- O'Hagan (1992).
- Ulman et al., 1993.
- Berns C.A. and Thomas C.A. (1965) J. Mol. Biol. **11**:476-490.
- Thomas W.R. and Rossi A.A. (1986) Infect. Immun. **52**:812-817.
- Thomas W.R. et al. (1990) Infect. and Immun. **58**:1090-1913.
- Caralone G.M. et al. (1986) J. Clin. Microbiol. **24**:330-331.
- Smith, D.B. and Johnson K.S. (1988) Gene **67**:31-40.
- Harkness, R. et al. (1992) J. Bacteriol. **174**:2425-2430.
- Hamel et al. (1987) J. Med. Microbiol. **23**:163-170.
- Mills et al. (1993) Infect. Immun. **61**:399-410.
- Trinchieri, (1993) Immunol. Today **14**:335-338.
- Hope, T.P. (1986) J. Immunol. Methods **88**:1-18.
- Zangwill et al., 1993. MMWR **42**:1-15.
- Loeb et al. 1987. Infect. Immun. **55**:2612-2618.
- Panezutti, 1993. Infect. Immun. **61**: 1867-1872.

CLAIMS

What we claim is:

1. A purified and isolated nucleic acid molecule, the molecule comprising at least a portion coding for a D15 outer membrane protein and having a DNA sequence selected from (a) the DNA sequence set forth in any one of Figures 1A to 1E or its complementary strand, and (b) DNA sequences which hybridize under stringent conditions to the DNA sequences defined in (a).
2. The molecule of claim 1 wherein said DNA sequences defined (b) have at least 90% sequence identity with the sequences defined in (a).
3. The molecule of claim 1 wherein said DNA sequences defined in (b) comprise the consensus sequence set forth in Figure 1F.
4. A recombinant plasmid adapted for transformation of a host, the recombinant plasmid comprising a plasmid vector into which has been inserted a DNA segment comprising at least an 18 bp fragment of a DNA molecule of claim 1, 2 or 3
5. The recombinant plasmid of claim 4 which is selected from plasmid DS-712-2-1 having ATCC accession number 75604 deposited November 4, 1993 and plasmid JB-1042-5-1 having ATCC accession number 75606 deposited November 4, 1993.
6. A recombinant vector adapted for transformation of a host cell, the recombinant vector comprising at least a DNA segment comprising at least an 18 bp fragment of a DNA molecule of claim 1, 2 or 3 and expression means operatively coupled to the DNA segment for expression of the gene product encoded thereby in the host cell.
7. The recombinant vector of claim 6 being plasmid DS-880-1-2 having ATCC accession number 75605 deposited November 4, 1993 and encoding the D15 gene product of H. influenzae SB33.

8. The recombinant vector of claim 6 wherein said DNA segment encodes a polypeptide of at least 6 residues.
9. The recombinant vector of claim 8 wherein said polypeptide is selected from those shown in Table 2.
10. The recombinant vector of claim 6, 7, 8 or 9 wherein said DNA segment consists of no more than the coding sequence for said D15 outer membrane protein.
11. The recombinant vector of claim 10, wherein the DNA segment further comprises a nucleic acid sequence encoding a leader sequence for export of said gene product from said host.
12. A purified and isolated protein encoded by the DNA fragment contained in the recombinant vector of claim 10 or 11.
13. A purified and isolated D15 outer membrane protein, or a portion thereof.
14. The protein of claim 13 wherein the D15 outer membrane protein is a Haemophilus D15 outer membrane protein.
15. The protein of claim 14 wherein the D15 outer membrane protein is a Haemophilus influenzae D15 outer membrane protein.
16. The protein of claim 15 wherein the Haemophilus influenzae is a type b Haemophilus influenzae strain.
17. The protein of claim 16 wherein the Haemophilus influenzae type b strain is selected from Ca, MinnA and Eagan strains.
18. The protein of claim 15 wherein the Haemophilus influenzae is a non-typeable Haemophilus influenzae strain.
19. The protein of claim 18 wherein the non-typeable Haemophilus influenzae strain is selected from PAK12085 and SB33 strains.
20. A synthetic peptide containing an amino acid sequence corresponding to the amino acid sequence of the protein or portion thereof claimed in any one of claims

- 11 to 19, or variant or mutant which retains immunogenicity.
21. The peptide of claim 20, having an amino acid sequence selected from those contained in Table 2.
22. An immunogenic composition, comprising a nucleic acid molecule claimed in any one of claims 1 to 5, a protein as claimed in any one of claims 12 to 14 or a peptide as claimed in claim 20 or 21 and a physiologically-acceptable carrier therefor.
23. The immunogenic composition of claim 22 formulated as a vaccine for in vivo administration to protect against diseases caused by Haemophilus.
24. The immunogenic composition of claim 23 formulated as a microparticle preparation, capsule preparation or liposome preparation.
25. The immunogenic composition of claim 23 in combination with a targeting molecule for delivery to specific cells of the immune system or to mucosal surfaces.
26. A method for inducing protection against disease caused by Haemophilus, comprising administering to a subject an effective amount of the immunogenic composition of any one of claims 22 to 25 to provide protective immunity against Haemophilus disease.
27. An antiserum or an antibody specific for a protein, a peptide or an immunogenic composition of any one of claims 12 to 25.
28. A chimeric molecule, comprising a protein as claimed in any one of claims 12 to 19 or a peptide claimed in claims 20 to 21 linked to another polypeptide or protein or a polysaccharide.
29. The chimeric molecule of claim 28 wherein said another polypeptide or protein comprise a surface protein or peptide corresponding thereto from a pathogenic bacteria.

30. The chimeric molecule of claim 29 wherein said another polypeptide or protein comprises a P1, P2 or P6 outer membrane protein of H. influenzae.

31. The chimeric molecule of claim 28 wherein said polysaccharide comprises a PRP molecule from H. influenzae.

FIG. 1A.

#### **H. influenzae b Ca strain D15 sequence**

## **SUBSTITUTE SHEET**

## FIG. 1A.(CONTINUED)

YS. ILE LYS GLY ASN SER VAL ILE PRO THR GLU ALA LEU LYS GLN ASN LEU ASP ALA ASN G  
AAA T C A A A G G T A A C T C T G T T A T T C C C A C T G A A G C A C T T A A C A A A A C T T A G C T A A C G  
370 390 400 410 420

LY PHE LYS VAL GLY ASP VAL LEU ILE ARG GLU LYS LEU ASN GLU PHE ALA LYS SER VAL L  
G G T T A A A G T T G G C G A T G T T A T T C G A G A A A A T T A A T G A A T T G C C A A A A G T G T A A  
430 440 450 460 470 480

YS. GLU HIS TYR ALA SER VAL GLY ARG THR ASN ALA THR VAL GLU PRO ILE VAL ASN THR L  
A A G A G C A C T A T G C A A G T G T A G G T C G C T A T A C G C A A C A G T T G A A C C T A T T G T C A A T A C G C  
490 500 510 520 530 540

EU PRO ASN ASN ARG ALA GLU ILE LEU ILE GLN ILE ASN GLU ASP ASP LYS ALA LYS LEU A 2/8 2  
T A C C A A A T A A T C G G G C T G A A A T T T A A T T C A A A T C A A T G A A G A T G A T A A A G C A A A A T T G G  
550 560 570 580 590 600

IA SER LEU THR PHE IYS GLY ASN GLU SER VAL SER SER THR LEU GLN GLN MET G  
C A T C A T T A A C T T C A A G G G G A A C G A A T C T G T T A G T A G C A G T A C A T T A C A A G A A C A A A T T G G  
610 620 630 640 650 660

IU LEU GLN PRO ASP SER TRP TRP IYS LEU TRP GLY ASN LYS PHE GLU GLN ALA GLN PHE G  
A A T T A C A A C C T G A T T C T T G G T G G A A A T T A T G G G G A A A T A A T T G A A G G T G C G C A A T T C G  
670 680 690 700 710 720

end truncated GST/D15

IU LYS ASP LEU GLN SER ILE ARG ASP TYR TYR LEU ASN ASN GLY TYR ALA LYS ALA GLN I  
A G A A G A T T G C A G T C A A T T C G T G A T T A T T A A T G G C T A T G C C A A A G C A C A A  
730 740 750 760 770 780

## FIG.1A.(CONTINUED)

LE	THR	LYS	THR	ASP	VAL	GLN	LEU	ASN	ASP	GLU	LYS	THR	LYS	VAL	ASN	VAL	THR	ILE	ASP	V
T T A C T A A A A C G G A T G T T C A G C T A A T G A T G A A A C A A A A C A A A G T T A A T G T A A C C A T T G A T G	800																			
AL	ASN	GLU	GLY	LEU	GLN	TYR	ASP	LEU	ARG	SER	ALA	ARG	ILE	IIE	GLY	ASN	LEU	GLY	GLY	
T A A T G A A G G T T A C A G T A T G A C C T T C G T A G T G C A C G C A T T A T A G G T A A T C T G G G A G G T A	850																			
ET	SER	ALA	GLU	LEU	GLU	PRO	LEU	LEU	SER	ALA	LEU	HIS	LEU	ASN	ASP	THR	PHE	ARG	ARG	S
T G T C T G C C G A G C T T G A A C C T T A C T T C A G C A T T A C A T T A A T G A T A C T T C C G C C G T A	900																			
ER	ASP	IIE	ALA	ASP	VAL	GLU	ASN	ALA	ILE	LYS	ALA	LYS	LEU	GLY	GLU	ARG	GLY	TYR	GLY	S
G T G A T A T T G C A G A T G T A G A A A T G C A A T T A A A G C A A A A C T T G G A G A C C G G T T A C G G T A	950																			
ER	ALA	THR	VAL	ASN	SER	VAL	PRO	ASP	PHE	ASP	ASP	ALA	ASN	LYS	THR	LEU	ALA	IIE	THR	L
G C G C A A C G G T A A A T T C A G T A C C T G A T T G C A A A T A A A C A T T A G C G A T A A C C C	1000																			
EU	VAL	VAL	ASP	ALA	GLY	ARG	LEU	THR	VAL	ARG	GLN	LEU	ARG	PHE	GLU	GLY	ASN	THR	V	
T T G T T G T T G A T G C T G G A C G T T A A C T G T T C G C C A A C T T C G C T T G A A A G G A A A T A C C G	1050																			
	1100																			
	1120																			
	1130																			
	1140																			

**FIG. 1A.(CONTINUED)**

AL SER ALA ASP SER THR LEU ARG GLN GLU MET ARG GLN GLU GLY THR TRP TYR ASN S  
 T T T C T G C T G A T A G C A C T T A C G G A A T G C G C C A A C A A G A A G G A A C T T G G T A T A T T 1200  
 1150 1160 1170 1180 1190  
  
 ER GLN LEU VAL GLU LEU GLY LYS ILE ARG LEU ASP ARG THR GLY PHE PHE GLU THR VAL G  
 C A C A A T T A G T T G A G T T A G G A A A A T T C G C T T A G A T C G T A C A G G T T C T T C G A A A C A G T C G 1260  
 1210 1220 1230 1240 1250  
  
 LU ASN ARG ILE ASP PRO ILE ASN GLY SER ASN ASP GLU VAL ASP VAL VAL TYR LYS VAL L  
 A A A A C C G A A T T G A T C C T A T C A A T G G T A G T A A T G A A G T G G A T G G A T G G T C G T A T A A A G T C A 1320  
 1270 1280 1290 1300 1310  
  
 YS GLU ARG ASN THR GLY SER ILE ASN PHE GLY ILE GLY TYR GLY THR GLU SER GLY ILE S  
 A A G A A C C G T A A C A C G G G T A G T A T C A A C T T G G T A T T G G T A C G G T A C A G A G A G T G G G T A T T A 1380  
 1330 1340 1350 1360 1370  
  
 ER TYR GIN ALA SER VAL LYS GLN ASP ASN PHE ILE GLY THR GLY TYR PRO TYR PHE T  
 G T T A T C A A G C A A G T G T T A A A C A A G A T A A T T C T T G G G A A C A G G G G G C A G T A A G T A T A G 1440  
 1390 1400 1410 1420 1430  
  
 LA GLY THR LYS ASN ASP TYR GLY THR SER VAL ASN LEU GLY TYR THR GLU PRO TYR PHE T  
 C T G G T A C G A A A A T G A T T A T G G T A C G A G T A C G A G T T G G G T T A T A C C G A G C C T A T T T A 1500  
 1450 1460 1470 1480 1490  
  
 HR LYS ASP GLY VAL SER LEU GLY GLY ASN VAL PHE PHE GLU ASN TYR ASP ASN SER LYS S  
 C T A A A G A T G G T G C T T G G A A A T G T T C T T G A A A A C T A C G A T A A C T C T A A A A 1560  
 1510 1520 1530 1540 1550

4/82

FIG. 1A. (CONTINUED)

FIG. 1A. (CONTINUED)

## FIG.1A.(CONTINUED)

ER THR GLY VAL GLY PHE GLN TRP GLN SER PRO ILE GLY PRO LEU VAL PHE SER TYR ALA L  
 C T A C A G G T G T C G G A T T C C A A T G G C A A T C T C T A T T G G C C A T T G G T A T T C T C T A T G C C A  
 2350 2360 2370 2380 2390 2400  
 YS PRO ILE LYS LYS TYR GLU ASN ASP ASP VAL GLU GLN PHE GLN PHE SER ILE GLY GLY S  
 A A C C A A T T A A A A A T A T G A A A A T G A T G A T G T C G A C A G T T C C A A T T A G T A T T G G A G G T T  
 2410 2420 2430 2440 2450 2460  
 ER PHE \*\*\* \*\*\*  
 C T T T C T A A T A A T T G A A C T T T C T T C A T C A G A A C T C A A A A C A A C G T T C T C T G C C T A A  
 2470 2480 2490 2500 2510 2520  
 T T T A A T T G G C A G A G A A A T A T T A A C C C A T C A T T A A G G A T A T T A T C A A T G A A  
 2530 2540 2550 2560 2570 2580  
 A A A C A T C G C A A A G T A A C C G C A C T T G C T T A G G T A T T G C A C T T G C T T C A G G C T A T G C T T C  
 2590 2600 2610 2620 2630 2640  
 C G C T G A A G A A A A T T G C T T C A T T A A T G C A G G T A T T G C A C T T C A A C A T C A C C C A G A T C G C  
 2650 2660 2670 2680 2690 2700  
 C A A G C G G T A G C A G A T A A A C T T G C T G C A T T G A G C T A G C T A G C A T T A G C A G C A  
 2710 2720 2730 2740 2750 2760

7/82

**FIG. 1A.(CONTINUED)**

AGC A A A A G A A G T T G A T G A T A A A A T T G C T G C T C G T A A A A A G T A G A A G C A A A A G T T  
2770 2780 2800 2810 2820

G C G G C T T A G A A A A G A T G C A C C T C G C T T A C G T C A A G C T G A T A T T C A A A A A C G C C A A C A G  
2830 2840 2850 2860 2870 2880

G A G A T T A A A A T T A G G T G C G G C T G A A G A T G C T G A A T T A C A A A A A T T A A T G C A A G A C A A  
2890 2900 2910 2920 2930 2940

G A T A A A A A A

8/82

**FIG. 1B.**

DS-712-2-1 DNA,  
Eagan D15 sequence  
IS THE SEQUENCE BEING TRANSLATED

A C A G G A C A G C T T C C C T T A A C C T T G A A A A T A T T A G G G A A A T T A C T T C C T G G C G A T T G 10 20 30 40 50 60	T C A T T A A T T A A G T G G G C C A A T T C T A T T G C A A A A G G T G C T G C C C A T C A G C A A A T 70 80 90 100 110 120	A T T G G A T T G G T G T A T T A A G T T T A A G T G C A C T G C A T G T G A T T A G T G T A A A T T A C G G A T T A T G 130 140 150 160 170 180	A A T T A T T C C A T T A C C A G T A T T A G A T G G C G G T C A T T A G T T T T T A A C A A T T G G G A A G C T 190 200 210 220 230 240	G T T A A A G G A A A C C T G T T C T G A G C G G G T G C A A A G C A T C T G T T A T C G A A T T G G C G C A G C A 250 260 270 280 290 300	C T G T T A T T A A G C T T A A C G G T G T T G C A T T A T T A A T G A T T T A C G T C T A T A A T T A T A 310 320 330 340 350 360
---	--	---	---	---	---

9/82

FIG. 1B. (CONTINUED)

TAGGATACAAATCGATGAAAACCTTCTAAATCGCAAGTTATTCCGGTACGACAAACGAC  
 370  
 MET LYS LYS LEU ILE ALA SER LEU LEU PHE GLY THR THR THR  
 TGTGTTGCCGCACCTTGTGGCAAAAGATAATTGGTGGATGGTGCCTGTTGACTGACCTT  
 430  
 VAL PHE AIA AIA PRO PHE VAL ALA LYS ASP ILE ARG VAL ASP GLY VAL GIN GLY ASP LEU  
 380  
 390  
 400  
 410  
 420  
 GLU GLN GIN ILE ARG ALA SER LEU PRO VAL ARG ALA GLY GIN ARG VAL THR ASP ASN ASP  
 AGAACAAATCCTGGCAAGTTACCTTGTTACCTGTTGGTGCCTGTTGACTGACAAATGAA  
 490  
 500  
 510  
 520  
 530  
 540  
 VAL ALA ASN ILE VAL ARG SER LEU PHE VAL SER GLY ARG PHE ASP ASP VAL LYS ALA HIS  
 TGTGGCTAAATTGTCCTGTTATTCGTAAGTAGGTGGTCGATTGATGAAAGGGCAA  
 550  
 560  
 570  
 580  
 590  
 600  
 GIN GLU GLY ASP VAL LEU VAL SER VAL ALA LYS SER ILE SER ASP VAL LYS  
 TCAAGAAAGGCGATGTCCTGTTAGGCTTGGCTTAAATCGATCATTTAGATGTTAA  
 610  
 620  
 630  
 640  
 650  
 660  
 ILE LYS GLY ASN SER VAL ILE PRO THR GLU ALA LEU LYS GIN ASN LEU ASP ALA ASN GLY  
 ATCAAAAGGTAAACTCTGTATTCCACTGAAGCACCTTAAACAAACTTAAACGG  
 670  
 680  
 690  
 700  
 710  
 720

## FIG.1B.(CONTINUED)

PHE	LYS	VAL	GLY	ASP	VAL	LEU	ILE	ARG	GLU	LYS	LEU	ASN	GLU	PHE	ALA	LYS	SER	VAL	LYS	
G T T A A G T T G G C G A T G T T A T T C G A G A A A A T T A A T G A A T T G C C C A A A A G T G T A A A	730	740	750	760	770	780														
GLU	HIS	TYR	ALA	SER	VAL	GLY	ARG	TYR	ASN	ALA	THR	VAL	GLU	PRO	ILE	VAL	ASN	THR	LEU	
A G A G C A C T A T G C A A G T G T G T A G G T C G C T A T A A C G C A A C A G T T G A A C C T A T T G T C A A T A C G C T	790	800	810	820	830	840														
PRO	ASN	ASN	ARG	ALA	GLU	ILE	LEU	ILE	GLN	ILE	ASN	GLU	ASP	ASP	LYS	ALA	LYS	LEU	ALA	
A C C A A A T A A T C G C G C T G A A A T T T A A T T C A A A T G A A G A T G A T A A A G C A A A A T T G G C	850	860	870	880	890	900														
SER	LEU	THR	PHE	LYS	GLY	ASN	GLU	SER	SER	SER	SER	SER	THR	LEU	GLN	GLU	GLN	MET	GLU	
A T C A T T A A C T T C A A G G G A A C G A A T C T G T A G T A G C A G T A C A T T A C A A G A A C C A A A T T G G A	910	920	930	940	950	960														
LEU	GLN	PRO	ASP	SER	TRP	TRP	LYS	LEU	TRP	GLY	ASN	LYS	PHE	GLU	GLY	ALA	GLN	PHE	GLU	
A T T A C A A C C T G A T T C T T G G T G G A A A T T A T G G G G A A A T T A A T G C C T A T G C C G C G C A A T T C G A	970	980	990	1000	1010	1020														
LYS	ASP	LEU	GIN	SER	ILE	ARG	ASP	TYR	TYR	LEU	ASN	GLY	TYR	ALA	LYS	ALA	GLN	ILE		
G A A A G A T T G C A G T C A A T T C G T G A T T A T T A A T G G C T A T G C C C A A A G C A C A A A T	1030	1040	1050	1060	1070	1080														
THR	LYS	THR	ASP	VAL	GIN	LEU	ASN	ASP	GLU	LYS	THR	LYS	VAL	ASN	VAL	THR	ILE	ASP	VAL	
T A C T A A A C G G A T G T T C A G C T A A T G A T G A A A A C A A A A G T T A A T G T A A C C A T T G A T G T	1090	1100	1110	1120	1130	1140														

FIG. 1B. (CONTINUED)

FIG.1B.(CONTINUED)

FIG. 1B. (CONTINUED)

## FIG.1B.(CONTINUED)

ARG	LEU	PRO	PHE	TYR	GLN	THR	TYR	THR	ALA	GLY	GLY	IIE	GLY	SER	LEU	ARG	GLY	PHE	ALA	
G C G T T A C C G T T C T A T C A A C T A C A G C G G C A T C G G T T C A T T A C G T G G T T G G T T G C	2290																			
																				2340
TYR	GLY	SER	IIE	GLY	PRO	ASN	ALA	IIE	TYR	ALA	GLU	TYR	GLY	ASN	GLY	SER	GLY	THR	GLY	
T T A T G G T A G T T G G A C C T A A C G C A A T T T A T G C C G A A T A T G G T A A T G G T A G T G G T A C T G G	2350																			
																				2390
THR	PHE	LYS	LYS	IIE	SER	SER	ASP	VAL	IIE	GLY	GLY	ASN	ALA	IIE	ALA	THR	ALA	SER	ALA	
T A C T T T A A G A A G A T A A G T T C T G A T T G T G A T T G G T G G T A A T G C A A T C G C T A C A G C T A G C G G C	2360																			
																				2400
THR	PHE	LYS	LYS	IIE	SER	SER	ASP	VAL	IIE	GLY	GLY	ASN	ALA	IIE	ALA	THR	ALA	SER	ALA	
T A C T T T A A G A A G A T A A G T T C T G A T T G T G A T T G G T G G T A A T G C A A T C G C T A C A G C T A G C G G C	2410																			
																				2450
GLU	LEU	IIE	VAL	PRO	THR	PRO	PHE	VAL	SER	ASP	LYS	SER	GLN	ASN	THR	VAL	ARG	THR	SER	
A G A G T T A T T G T G C C A A C T C C A T T G T G A G C G A T A A G A G C C A A A A T A C G G T C C G A A C C T C	2420																			
																				2460
GLU	LEU	IIE	VAL	PRO	THR	PRO	PHE	VAL	SER	ASP	LYS	SER	GLN	ASN	THR	VAL	ARG	THR	SER	
A G A G T T A T T G T G C C A A C T C C A T T G T G A G C G A T A A G A G C C A A A A T A C G G T C C G A A C C T C	2470																			
																				2510
LEU	PHE	VAL	ASP	ALA	ALA	SER	VAL	TRP	ASN	THR	LYS	TRP	LYS	SER	ASP	LYS	ASN	GLY	LEU	
C T T A T T G T T G A T G C G G C A A G T G T T G G A A T A C T A A A T G G A A A T C A G A T A A A A T G G A T T	2530																			
																				2570
GLU	SER	ASP	VAL	LEU	LYS	ARG	LEU	PRO	ASP	TYR	GLY	LYS	SER	SER	ARG	IIE	ARG	ALA	SER	
A G A G A G C G G A T G T A T T A A A A G A T T G C C T G A T T G C C A A A T C A A G C C G C T A T T C G C G C T C	2590																			
																				2630
																				2640

## FIG.1B.(CONTINUED)

THR	GLY	VAL	GLY	PHE	GIN	TRP	GIN	SER	PRO	ILE	GLY	PRO	LEU	PHE	SER	TYR	ALA	LYS
T A C A G G T G T C G G A T T C C A A T G G C A A T C T C C T A T T G G G C A T T G G T A T T C T C T A T G C C A A																		
2650	2660	2670	2680	2690	2700	2710	2720	2730	2740	2750	2760	2770	2780	2790	2800	2810	2820	16/82
PRO	ILE	LYS	LYS	TYR	GLU	ASN	ASP	VAL	GLU	GLN	PHE	GLN	PHE	SER	ILE	GLY	GLY	SER
A C C A A T T A A A A A T G A A A A T G A T G A T G A C A G T T C C A A T T A G T A T G A C A G T C G A A T G T C																		
2710	2720	2730	2740	2750	2760	2770	2780	2790	2800	2810	2820	2830	2840	2850	2860	2870	2880	2890
PHE	***	***																
T T T C T A A T A A T T G A A C T T T C T T C A T C A T C A T C A A C G T T C T C T G C C T A A T																		
2770	2780	2790	2800	2810	2820	2830	2840	2850	2860	2870	2880	2890	2900	2910	2920	2930	2940	16/82
T T A A T T G G C A G A G A A A T T A A A C C C A T C A T T A A G G A T A T T C A A A T G C A A																		
2830	2840	2850	2860	2870	2880	2890	2900	2910	2920	2930	2940	2950	2960	2970	2980	2990	2990	16/82
A A C A T C G C A A A G T A A C C G C A C T T G C T T A G G T A T G C A C T T G C T A T G C T T C C																		
2890	2900	2910	2920	2930	2940	2950	2960	2970	2980	2990	2990	2990	2990	2990	2990	2990	2990	16/82
G C T G A A G A A A A A T T G C T T C A T T A A T G C A C T T A T T C A A																		
2950	2960	2970	2980	2990	2990	2990	2990	2990	2990	2990	2990	2990	2990	2990	2990	2990	2990	16/82

## FIG.1C.

DS-691-1-5 DNA, M1m A D15 sequence  
IS THE SEQUENCE BEING TRANSLATED

A A T C A C T T A C T G G C A T T G T C A T T A A T A A T T A A G T G G G C C A A T T G C A A A A G  
10 20 30 40 50

G T G C T G G C A C A T C A G C A A A T A T T G G A T T G G T G T A T T A A G T T T A A G T T T A A G T G G C A C T G A T T A  
70 80 90 100 110 120  
17/82

G T G T A A A T T T A G G G A T T A T T G A A T T G A A T T C C A T T A C C A G T A T T A G A T G G C G G T C A T T T A G  
130 140 150 160 170 180

T T T T T T A A C A A T G G A A G C T G T T A A A G G A A A C C T G T T C T G A G C G G G T G C A A A G C A T C T  
190 200 210 220 230 240

G T T A T C G A A T T G G G C A G G C A C T G T T A A G C T T A A G C T T A A C G G T G T G C A T T A T G A T T  
250 260 270 280 290 300

## FIG. 1C.(CONTINUED)

T T T T A C G T C T A T T A T T A G G A T A C A A T C G A T G A A A A A C T T C T A A T C G C A A G T T A  
 310 320 330 340 350 360  
 MET LYS LYS LEU ILE ALA SER LEU  
  
 LEU PHE GLY THR THR VAL PHE ALA ALA PRO PHE VAL ALA LYS ASP ILE ARG VAL  
 T T A T T C G G T A C G A C A A C G A C T G T G T T G C C G C A C C T T G T G T G C A A A G A T A T T C G T G T G  
 370 380 390 400 410 420  
  
 ASP GLY VAL GLN GLY ASP LEU GLU GIN GIN ILE ARG ALA SER LEU PRO VAL ARG ALA GLY  
 G A T G G T G T C A A G G T G A C T T A G A A C A A C A A T C C G A G C A A G T T A C C T G T T C G T G C C G G T  
 430 440 450 460 470 480 490  
 18 / 82  
  
 GIN ARG VAL THR ASP ASN ASP VAL ALA ASN ILE VAL ARG SER LEU PHE VAL SER GLY ARG  
 C A G C G T G T G A C T G A C A A T G A T G T G C T A A T A T T G T C C G C T C T C G T A A G T G G T C G A  
 500 510 520 530 540  
  
 PHE ASP ASP VAL LYS ALA HIS GIN GLU GLY ASP VAL LEU VAL SER VAL VAL ALA LYS  
 T T C G A T G A T G T G A A A G C G C A T C A A G G A A G G C A T G T G C T T G T G C T A A A  
 550 560 570 580 590 600  
  
 SER ILE ILE SER ASP VAL LYS ILE LYS GLY ASN SER VAL ILE PRO THR GLU ALA LEU LYS  
 T C G A T C A T T C A G A T G T T A A A A T C A A A G G T A A C T C T G T T A T T C C C A C T G A A G C A C T T A A A  
 610 620 630 640 650 660

FIG. 1C. (CONTINUED)

## FIG.1C.(CONTINUED)

GLY TYR ALA LYS ALA GIN ILE THR LYS THR ASP VAL GIN LEU ASN ASP GLU LYS THR LYS  
 G G C T A T G C C A A A G C A C A A T T A C T A A A C G G A T G T T C A G C T A A T G A T G A A A A C A A A A  
 1030 1040 1050 1060 1070 1080

VAL ASN VAL THR ILE ASP VAL ASN GLU GLY LEU GIN TYR ASP LEU ARG SER ALA ARG ILE  
 G T T A A T G T A A C C A T T G A T G T A A T G A A G G T T A C A G T A C A G T A C C T T C G T A G T G C A C G C A T T  
 1090 1100 1110 1120 1130 1140

ILE GLY ASN LEU GLY GLY MET SER ALA GLU LEU GLU PRO LEU LEU SER ALA LEU HIS LEU  
 A T A G G T A A T C T G G A G G T A T G T C T G C C G A G C T T A C T T C A G C A T T A C A T T A  
 1150 1160 1170 1180 1190 1200

O/82

ASN ASP THR PHE ARG ARG SER ASP ILE ALA ASP VAL GLU ASN ALA ILE LYS ALA LYS LEU  
 A A T G A T A C T T T C C G C C G T A G T G A T A T T G C A G A T G T A G A A A A T G C A A T T A A A G C A A A A C T T  
 1210 1220 1230 1240 1250 1260

GLY GLU ARG GLY TYR GLY SER ALA THR VAL ASN SER VAL PRO ASP PHE ASP ASP ALA ASN  
 G G A G A A C G C G G T T A C G G T A G C G C A A C G G T A A A T T C A G T A C C T G A T T G A T G C A A A T  
 1270 1280 1290 1300 1310 1320

LYS THR LEU ALA ILE THR LEU VAL VAL ASP ALA GLY ARG ARG LEU THR VAL ARG GIN LEU  
 A A A A C A T T A G C G A T A A C C C T T G T T G A T G C T G G A C G A C G T T A A C T G T T C G C C A A C T T  
 1330 1340 1350 1360 1370 1380

FIG. 1C. (CONTINUED)

## **SUBSTITUTE SHEET**

## FIG.1C.(CONTINUED)

TYR THR GLU PRO TYR PHE THR LYS ASP GLY VAL SER LEU GLY ASN VAL PHE PHE GLU  
 T A T A C C G A G C C C T A T T T A C T A A A G A T G G T G T A A G T C T T G G A A A T G T T C T T G A A  
 1750 1760 1770 1780 1790 1800

ASN TYR ASP ASN SER LYS SER ASP THR SER ASN TYR LYS ARG THR TYR GLY SER  
 A A C T A C G A T T A A C T C T A A A G T G A T A C A T C C T C T A A C T A T A A G C G T A C G A C T T A C C G A A G T  
 1810 1820 1830 1840 1850 1860

ASN VAL THR LEU GLY PHE PRO VAL ASN GLU ASN ASN SER TYR TYR VAL GLY LEU GLY HIS  
 A A T G T T A C T T A G G T T T C C T G T A A A T G A A A T A A C T C C T A T T G T A G G T A G G T C A T  
 1870 1880 1890 1900 1910 1920

THR TYR ASN LYS ILE SER ASN PHE ALA LEU GLU TYR ASN ARG ASN LEU TYR ILE GLN SER  
 A C C T A T A A A T T A G T A A C T T G C T G A A T A A C C G T A T T A A C C G T A T T A A T C A A T C A  
 1930 1940 1950 1960 1970 1980

MET LYS PHE LYS GLY ASN GLY ILE LYS THR ASN ASP PHE ASP SER PHE GLY TRP ASN  
 A T G A A A T T A A A G G T A A T G G C A T T A A A C A A T G A C T T G A T T T C T T T G G T T G G A A C  
 1990 2000 2010 2020 2030 2040

TYR ASN SER LEU ASN ARG GLY TYR PHE PRO THR LYS GLY VAL LYS ALA SER LEU GLY GLY  
 T A T A A C A G C C T T A A T A G A G G C T A T T C C A A C T A A A G G G T T A A A G C A A G T C T T G G T G G A  
 2050 2060 2070 2080 2090 2100

SUBSTITUTE SHEET

22/82

**FIG.1C.(CONTINUED)**

ARG VAL THR ILE PRO GLY SER ASP ASN LYS TYR TYR LYS LEU SER ALA ASP VAL GLN GLY  
 C G A G T T A C T A T T C C C A G G T T C T G A T A C A A A T A C T A C A A C T A A G T G C A G A T G T A C A G G G T  
 2110 2120 2130 2140 2150 2160  
 PHE TYR PRO LEU ASP ARG ASP HIS ILE TRP VAL VAL SER ALA LYS ALA SER ALA GLY TYR  
 T T C T A C C C A T T A G A C A G A G A T C A C C T C T G G G T T G T A T C T G C A A A G C A T C T G C A G G A T A T  
 2170 2180 2190 2200 2210 2220  
 23/82  
 ALA ASN GLY PHE GLY ASN LYS ARG ILE PRO PHE TYR GLN THR TYR THR ALA GLY GLY ILE  
 G C A A A T G G T T T G G A A C A A G C G T T A C C G T T C T A T C A A A C T T A T A C A G C G G T G G C A T C  
 2230 2240 2250 2260 2270 2280  
 GLY SER LEU ARG GLY PHE ALA TYR GLY SER ILE GLY PRO ASN ALA ILE TYR ALA GLU TYR  
 G G T T C A T T A C G T G G T T G C T T A T G G T A G T A T T G G A C C T A A C G C A A T T A T G C C G A A T A T  
 2290 2300 2310 2320 2330 2340  
 GLY ASN GLY SER GLY THR GLY THR PHE LYS LYS ILE SER SER ASP VAL ILE GLY GLY ASN  
 G G T A A T G G T A G T G G T A C T T G G T A C T T A A G A G A T A A G T G A T G T C T G A T G T G G T A A T  
 2350 2360 2370 2380 2390 2400  
 ALA ILE ALA THR ALA SER ALA GLU LEU ILE VAL PRO THR PRO PHE VAL SER ASP LYS SER  
 G C A A T C G C T A C A G C T A G C G C A G A G T T A A T T G T G C C A A C T C C A T T T G T G A G C G A T A A G A G C  
 2410 2420 2430 2440 2450 2460

## FIG.1C.(CONTINUED)

2470

GLN	ASN	THR	VAL	ARG	THR	SER	LEU	PHE	VAL	ASP	ALA	ALA	SER	VAL	TRP	ASN	THR	LYS	TRP	
C	A	A	A	T	A	C	G	G	T	C	C	G	A	C	C	T	T	G	T	T

2480

AA	A	T	C	A	G	A	T	A	G	A	G	A	T	G	T	A	T	G	G
----	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

2490

2500	2510
------	------

2530

LYS	SER	ASP	LYS	ASN	GLY	LEU	GLU	SER	ASP	VAL	LEU	LYS	ARG	LEU	PRO	ASP	TYR	GLY	LYS
A	A	T	C	A	G	A	T	A	G	A	G	A	T	G	T	G	C	T	G

2540

2550	2560
------	------

2590

SER	SER	ARG	ILE	ARG	ALA	SER	THR	GLY	VAL	GLY	PHE	GLN	TRP	GLN	SER	PRO	ILE	GLY	PRO
T	C	A	A	G	C	C	G	T	A	C	A	G	G	A	T	G	C	A	T

2600

2610	2620
------	------

2650

LEU	VAL	PHE	SER	TYR	ALA	LYS	PRO	ILE	LYS	TYR	GLU	ASN	ASP	ASP	VAL	GLU	GLN	PHE	
T	T	G	G	T	A	T	C	T	T	A	T	G	C	C	T	C	G	A	T

2660

2670	2680
------	------

2710

GLN	PHE	SER	ILE	GLY	GLY	SER	PHE	***	***	***	***	***	***	***	***	***	***	***	
C	A	A	T	T	A	G	T	A	T	G	G	G	T	T	C	T	T	C	A

2720

2730	2740
------	------

2750

2760	2770
------	------

## FIG.1C.(CONTINUED)

A A C A C G T T C T C T G C C T A A T T A A T T G G G C A G A G A A A T T A A C C C A T T C A T T A A T T A  
2770 2780 2800 2810 2820

A G G A T A T T A T C A A A A T G A A A A C A T C G C A A A G T A A C C G C A C T T G C T T A G G T A T T G C A C  
2830 2840 2850 2860 2880

T T G C T T C A G G C T A T G C T T C C G C T G A A G A A A A A T T G C T T C A T T A A T G C G G G T A T A N T T 25/82  
2890 2900 2910 2920 2930 2940

T N C A A G G C N A A G G  
2950

## FIG. 1D.

SB3 D15

IS THE SEQUENCE BEING TRANSLATED

G G C A T T G A A A A A C A G G A C A G C T T C C C T T A A C C T T G A A A A T A T T A G G G A A A T T A C C T T  
10 20 30 40 50 60

A C T G G C G A T T T G T C A T T A A T T A A G T G G G C C A A T T T C T A T T G C A A A A G G T G C T G G C  
70 80 90 100 110 120

26/82  
G C A T C A G C A A A T A T T G G A T T G G T G T A T T G G C A T T G A T T G T G T A A A T T  
130 140 150 160 170 180

T T A G G G A T T A T G A A T T T A T T C C A T T A C C A G T A T T A G G A T T G G C G G T C A T T A G T T T T T A  
190 200 210 220 230 240

A C A A T G G A A G G C T G T T A A A G G A A A A C C T G T T C T G A G G C G G G T G C A A A A G C A T C T G T T A T C G A  
250 260 270 280 290 300

FIG. 1D. (CONTINUED)

## FIG. 1D.(CONTINUED)

SER	ASP	VAL	LYS	ILE	LYS	GLY	ASN	SER	ILE	ILE	PRO	PRO	GLU	ALA	LEU	LYS	GLN	ASN	LEU
T T C A G A T G T T A A A T C A A A G G T A A C T C T A T T A T T C C T G A A G C A C T A A A C A A A C T T	670																		
680																			720
ASP	ALA	ASN	GLY	PHE	LYS	VAL	GLY	ASP	ILE	LEU	ILE	ARG	GLU	LYS	LEU	ASN	GLU	PHE	ALA
A G A T G C T A A C G G G T T A A A G T T G G C G A T A T T C G A G A A A T T A A A T G A A T T G C	730																		
740																			780
GLN	SER	VAL	LYS	GLU	HIS	TYR	ALA	SER	VAL	GLY	ARG	TYR	ASN	ALA	THR	VAL	GLU	PRO	ILE
C C A A G T G T A A A G A G C A C T A T G C A A G T G T A G G T C G C T A T A A C G C A A C C G T T G A A C C T A T	790																		830
800																			840
VAL	ASN	THR	LEU	PRO	ASN	ARG	ALA	GLU	ILE	LEU	ILE	GLN	ILE	ASN	GLU	ASP	ASP	LYS	28 / 82
T G T C A A T A C G C T A C C A A A T A A T C G C G C T G A A A T T T A A T T C A A A T C A A T G A A G A T G A T A A	850																		
860																			
VAL	ASN	THR	LEU	PRO	ASN	ARG	ALA	GLU	ILE	LEU	ILE	GLN	ILE	ASN	GLU	ASP	ASP	LYS	
910																			
920																			
ALA	LYS	LEU	ALA	SER	LEU	THR	PHE	LYS	GLY	ASN	GLU	SER	VAL	SER	SER	SER	THR	LEU	GIN
A G C C A A A T T G G C A T C A T T A A C T T C A A C C T G A T T G G G A A C G A A T C T G T T A G C A G T A C A T T A C A	930																		
940																			
950																			
960																			
GLU	GLN	MET	GLU	ILE	GLN	PRO	ASP	SER	TRP	TRP	LYS	LEU	TRP	GLY	ASN	LYS	PHE	GLU	GLY
A G A A C A A A T T G G A A T T A C A A C C T G A T T G G G A A A T T A T G G G G A A T A A T T G A A G G	970																		
980																			
1000																			
1010																			
1020																			

## FIG.1D.(CONTINUED)

ALA	GLN	PHE	GLU	LYS	ASP	LEU	GIN	AIA	ILE	ARG	ASP	TYR	TYR	LEU	ASN	ASN	GLY	TYR	ALA	
T G C G C A A T T C G A G A A A G A T T G C A G G C A A T T C G T G A T T A T T A A T A T G G C T A T G C																				
1030																				1080
LYS	AIA	GIN	ILE	THR	LYS	ALA	ASP	VAL	GIN	LEU	ASN	ASP	GLU	LYS	THR	LYS	VAL	ASN	VAL	
C A A G C A C A A A T C A C T A A A G C G A T G T T C A G C T A A A T G A T G A A A A A C A A A A G T T A A T G T																				
1090																				1140
THR	ILE	ASP	VAL	ASN	GLU	GLY	LEU	GIN	TYR	ASP	LEU	ARG	SER	ALA	ARG	ILE	ILE	GLY	ASN	
A A C C A T T G A T G T A A A T G A A G G T T A A C G T A T G A C C T T C G T A G T G C A C G C A T T A G G T A A																				
1150																				1200
LEU	GLY	GLY	MET	SER	ALA	GLU	LEU	GLU	PRO	LEU	LEU	SER	ALA	LEU	HIS	LEU	ASN	ASP	THR	
T C T G G G A G G T A T G T C T G C C G A G C T T G A A C C T T A C T T C A G C A T T A C A T T A A T G A T A C																				
1210																				1260
PHE	ARG	ARG	SER	ASP	ILE	ALA	ASP	VAL	GLU	ASN	ALA	ILE	LYS	LEU	GLY	GLU	ARG			
T T T C C G C C G T A G T G A T A T T G C A G T A G T G A A A A T G C A A T T A A A G C A A A A C T T G G G G A A C G																				
1270																				1320
GLY	TYR	GLY	ASN	THR	THR	VAL	ASN	SER	VAL	PRO	ASP	PHE	ASP	ASP	ALA	ASN	LYS	THR	LEU	
A G G T T A C G G T A A C A C A C A C A G T A A A T T C T G T A C C T G A T T G A C G A T T G C A A A T A A A A C A T T																				
1330																				1380

## FIG.1D.(CONTINUED)

AIA ILE THR PHE VAL ASP ALA GLY ARG ARG LEU THR VAL HIS GLN LEU ARG PHE PHE GLU  
 AGCGATAACCTTGTGTTGATGCTGGACGGAACTGTTAACCAACTTCCGCTTGA  
 1390 1400 1410 1420 1430 1440

GLY ASN THR VAL SER ALA ASP SER THR LEU ARG GLN GLU MET ARG GLN GLN GLU GLY GLY THR  
 AGGAAATAACCGTTCTGCTGATAGTAGTACCTTACGTCAGGAAATGCCAACAGAAC  
 1450 1460 1470 1480 1490 1500

TRP TYR ASN SER GIN LEU VAL GLU LEU GLY LYS ILE ARG LEU ASP ARG THR GLY PHE PHE  
 T TGGTATAATTCAAAATAGTTGAGTTAGGAAATACTCGCTTAGATCGTACAGGTT  
 1510 1520 1530 1540 1550 1560

GLU THR VAL GLU ASN ARG ILE ASP PRO ILE ASN GLY SER ASN ASP GLU VAL ASP VAL VAL  
 CGAACACAGTTGAAACCGAAATTGATCCTATCACATTGGTAGCCATGATGAAAGTGG  
 1570 1580 1590 1600 1610 1620

TYR IYS VAL LYS GLU ARG ASN THR GLY SER ILE ASN PHE GLY ILE GLY TYR GLY THR GLU  
 ATATAAAAGTCAAAAGAACGTAACAGTGGTAGTATCAACTTGTATTTCTTGGTTACAGA  
 1630 1640 1650 1660 1670 1680

SER GLY ILE SER TYR GIN ALA SER VAL LYS GLN ASP ASN PHE LEU GLY THR GLY ALA ALA  
 GAGTGGTATAGTTATCAAAAGCAAGTGTCAAAACAGATAATTCTTGGTTACGG  
 1690 1700 1710 1720 1730 1740

FIG. 1D. (CONTINUED)

## **SUBSTITUTE SHEET**

## FIG.1D.(CONTINUED)

LEU ASN ARG GLY TYR PHE PRO THR LYS GLY VAL LYS ALA SER LEU GLY GLY ARG VAL THR  
 C C T T A A T A G A G G C T A T T C C A A C T A A A G G G T T A A G C C A A G T C T T G G T G G A C G A G T T A C  
 2110 2120 2130 2140 2150 2160  
 ILE PRO GLY SER ASP ASN LYS TYR TYR LYS LEU SER ALA ASP VAL GLN GLY PHE TYR PRO  
 A T T C C A G G T T C T G A T A A C A A T A C T A C A A C T A A G T G C A G A T G T A C A G G G T T C T A C C C  
 2170 2180 2190 2200 2210 2220  
 LEU ASP ARG ASP HIS LEU TRP VAL VAL SER ALA LYS ALA SER ALA GLY TYR ALA ASN GLY  
 A T T A G A C A G A G A T C A C C T C T G G G T T G T A T C T G C A A A G C A T C T G C A G G A T A T G C A A A T G G  
 2230 2240 2250 2260 2270 2280  
 PHE GLY ASN LYS ARG LEU PRO PHE TYR THR THR ALA GLY GLY ILE GLY SER LEU  
 T T T G G A A A C A A G C G T T A C C G T T A C T A A A C T T A C A G C G G G T G G C A T T G G T T C A T T  
 2290 2300 2310 2320 2330 2340  
 ARG GLY PHE ALA TYR GLY SER ILE GLY PRO ASN ALA ILE TYR GLN GLN ASN ASN LYS  
 A C G C G G T T T G C T T A T G G T A G C A T T G G C C T A A C G C A A T T A T C A A G G T C A A A T A A T A A  
 2350 2360 2370 2380 2390 2400  
 PHE ASN LYS ILE SER SER ASP VAL ILE GLY GLY ASN ALA ILE ALA THR ALA SER ALA GLU  
 A T T T A A T A A G A T A A G T T C T G A T T G G T G A T T G G C C T A T G C C A A T C G C T A C A G C T A G C G C A G A  
 2410 2420 2430 2440 2450 2460

## FIG.1D.(CONTINUED)

LEU	ILE	VAL	PRO	THR	PRO	PHE	VAL	SER	ASP	LYS	SER	GLN	ASN	THR	VAL	ARG	THR	SER	LEU
G T T A A T T G C C A A C T C C A T T T G T G A G T A A G A G T C A A A T A C A G T C C G A A C C T C C C T	2470	2480	2490	2500	2510	2520													
PHE	VAL	ASP	ALA	ALA	SER	VAL	TRP	ASN	THR	LYS	TRP	LYS	SER	ASP	LYS	ASN	GLY	LEU	GLU
A T T T G T G A T G C G G C A A G T G T T G G A A T A C T A A A T G G A A T C A G A T A A A A T G G A T T A G A	2530	2540	2550	2560	2570	2580													
SER	ASN	VAL	LEU	LYS	ASP	LEU	PRO	ASP	TYR	GLY	LYS	SER	SER	ARG	THR	ARG	ALA	SER	THR
G A G C A A T G T C T T G A A A G A C T T A C C C G A T T A T G G C A A A T C A A G C C G T A C T C G C G C T C T A C	2590	2600	2610	2620	2630	2640													
GLY	VAL	GLY	PHE	GLN	TRP	GLN	SER	PRO	SER	GLY	PRO	VAL	VAL	PHE	SER	TYR	ALA	LYS	PRO
A G G T G T C G G A T T C C A A T G G C A A T C T C C T A G T G G A C C A G T G G T A T T T C T T A T G C T A A C C	2650	2660	2670	2680	2690	2700													
ILE	LYS	LYS	TYR	GLU	ASN	ASP	VAL	GLU	GLN	PHE	GLN	PHE	SER	ILE	GLY	GLY	SER	PHE	
A A T T A A A A A T G A A A A T G A T G A T G T C G G A A C A G T T C C A A T T A G T A T T G G G G T T C T T	2710	2720	2730	2740	2750	2760													
***      ***																			
C T A A T A A T T G A A C T T T C G T C A T C A G A A C A A A C A A C G T T C T C T G C C T A A T T A	2770	2780	2790	2800	2810	2820													

33/82

## FIG.1D.(CONTINUED)

ATGGGAGAAATTAAACCATTCATTAAAGGATAATTCAAATGAAAAC  
2830 2840 2850 2860 2870 2880

ATCGCCAAAGTAAACCGCACTTGCTTAGGTATGCCACTTGCACCTGGCTATGCCAGCT  
2890 2900 2910 2920 2930 2940

GAGAAAATTGCTTTATTAATGCCAGGTATA  
2950 2960 2970

34/82

**FIG. 1E.**

JB-1042-9-4 DNA, PAK D15  
IS THE SEQUENCE BEING TRANSLATED

A A A G G C A T T G A A A A A C A G G A C A A C T T C C C T T A A C C T T G A A A A T T A G G G A A T T  
10 20 30 40 50 60

A C T T A C T G G C G A T T T G T C A T T A A T T A A G T G G C C A A T T C T A T T G C A A A A G G T G C  
70 80 90 100 110 120

T G G T G C A T C A G C A A A T A T T G G A T T G G T G T A T T A A G T G G G C A T T G C A T T A G T G T  
130 140 150 160 170 180

A A A T T A G G G A T T A T G A A T T G A T T A T T C C A T T A C C A G T A T T A G A T G G C G G T C A T T A G T T  
190 200 210 220 230 240

T T T A C A A T T G G A A G C T G T T A A A G G A A A A C C T G T T C T G A G C G G G T G C A T T A G T T  
250 260 270 280 290 300

T C G A A T T G G C G C A G C A C T G T T A A G C T T A A C G G T G T T G C A T T A T G A T T A G T T  
310 320 330 340 350 360

## FIG.1E.(CONTINUED)

A C G T C T A T A T T T A T A T A G G A T A C A A T C G A T G A A A A A C T T C T A A T C G C A A G T T A T T A T	MET	LYS	LYS	LEU	LEU	ILE	ALA	SER	LEU	LEU	P
370	390	400				410					
HE	GLY	THR	THR	VAL	PHE	ALA	ALA	PRO	PHE	VAL	
T C G G T A C G A C A A C G A C T G T G T T G C C G C A C C T T G T G T G G C A A A A G A T A T T C G T G G A T G	430	440	450	460	470						
LY	VAL	GLN	GLY	ASP	LEU	GLU	GLN	IIE	ARG	ALA	
G T G T T C A A G G T G A C T T A G A A C A A C A A A T C C G A G C A A G T T A C C T G T T C G T G G C T G G T C A G C	490	500	510	520	530						
RG	VAL	THR	ASP	ASN	ASP	VAL	ALA	ASN	IIE	ARG	
G T G T G A C T G A C A A T G A T G T G G C T A A T A T T G T C C G C T C T C T A T T C G T A A G T G G T C G A T T C G	550	560	570	580	590						
SP	ASP	VAL	LYS	ALA	HIS	GLN	GLU	ASP	VAL	SER	
A T G A T G T G A A A G C G C A T C A A G A A G G C G A T G T G C T T G T G T A G C G T T G T G G C T A A A T C G A	610	620	630	640	650						
LE	IIE	SER	ASP	VAL	LYS	IIE	LYS	GLY	ASN	SER	
T C A T T C A G A T G T A A A A T C A A A G G T A A C T C T G T T A T T C C C A C T G A A G C A C T T A A A C C A A A	670	680	690	700	710						
											36 / 82
											540

## FIG.1E.(CONTINUED)

SN	LEU	ASP	ALA	ASN	GLY	PHE	LYS	VAL	GLY	ASP	VAL	LEU	ILE	ARG	GLU	LYS	LEU	ASN	GLU	P
ACTT	A G A T	G C T A	A C G G	G T T A	A A G T	T G C G	A G T G	T T A A T	T C G A	G A A A	A T T A A T	G A A T								
730	740	750	760	770	780															
HE	ALA	LYS	SER	VAL	LYS	GLU	HIS	TYR	ALA	SER	VAL	GLY	ARG	TYR	ASN	ALA	THR	VAL	GLU	P
T T G C	C A A A	A G T G	T A A A	G A G C	A C T A	T G C A	A G C A	A G T G	T A G C	A A G T	G C T A	T C G C	T G G T	A G G T	G T A G G	C A A C	C G T T	G A A C		
790	800	810	820	830	840															
RO	ILE	VAL	ASN	THR	LEU	PRO	ASN	ASN	ARG	ALA	GLU	ILE	LEU	ILE	GLN	IIE	ASN	GLU	ASP	A
C T A T	T G T C	A A T A	C G C T	G C A A	A T A T	C G T G	C T G A	A A T T	A T T A	A T T C	A A T T C	A A T T C	A A T T C	A A T T C	A A T T C	A A T T C	A A T T C	A A T T C	A A T T C	
850	860	870	880	890	900															
SP	LYS	ALA	LYS	LEU	ALA	SER	LEU	THR	PHE	LYS	GLY	ASN	GLU	SER	VAL	SER	SER	SER	THR	L
A T A A	A G C A	A A A T	T G G C	A T C A	T T A A	C T T A	A C T T	A C T T	C A A G	G G G A	A C G A	A T C T	G T T A	G T A G	T A G T A	G C A G	T A C A T	N	37/82	
910	920	930	940	950	960															
EU	GIN	GLU	GIN	MET	GLU	LEU	GIN	PRO	ASP	SER	TRP	TRP	LYS	LEU	TRP	GLY	ASN	LYS	PHE	G
T A C A	A G A A	C A A A	T G G A	A T T A	C A A C	T G A T	T G G T	T G G T	G A A A	T T A T	T G G G	G A A A	T T A T	T G G G	G A A A	T A A T	T G G C	T		
970	980	990	990	980	970															
IU	GLY	ALA	GIN	PHE	GLU	LYS	ASP	LEU	GIN	ALA	ILE	ARG	ASP	TYR	TYR	LEU	ASN	GLY	T	
A A G G	T G C G	C A A T	T C G A	G A A A	G A T C	T G C A	G G C A	A T T C	G T G A	T T A T	T G G T	T G G T	A T T C	T G G T	T G G T	A T T C	A T T C	A T T C	A T T C	
1030	1040	1050	1060	1070	1080															
YR	ALA	LYS	ALA	GIN	ILE	THR	LYS	THR	ASP	VAL	GIN	ILE	ASN	ASP	GLU	LYS	THR	LYS	VAL	A
A T G C	C A A A	G C A C	A A A T	C A C T	A A A T	C A C T	A A A C	A A A C	A C G G	A T G G	T C A G	C T A G	C T A G	C T A G	C T A G	C T A G	C T A G	C T A G	C T A G	
1090	1100	1110	1120	1130	1140															

## FIG. 1E. (CONTINUED)

SN	VAL	THR	ILE	ASP	VAL	ASN	GLU	GLY	LEU	GLN	TYR	ASP	LEU	ARG	SER	ALA	ARG	ILE	ILE	G
ATGTAACCATTGATGTAATGAAAGGTACAGTATGACCTTGCTAGTCGCTTACATTATAG																				
1150																				1190
LY	ASN	LEU	GLY	GLY	MET	SER	ALA	GLU	LEU	GLU	PRO	LEU	LEU	SER	ALA	LEU	HIS	LEU	ASN	A
GTAATCTGGAGGGTATGTCATGCTGCCGAGCTTGAAACCTTACTTACATTAAATAG																				
1210																				1250
SP	THR	PHE	ARG	ARG	SER	ASP	ILE	ALA	ASP	VAL	GLU	ASN	ALA	ILE	LYS	ALA	LYS	LEU	GLY	G
ATACTTTCCGCCGCTAGTAGTGTATGCAAGATATTGCAATTGCAAATTAAATGCAAAAGC																				38
1270																				1320
IU	ARG	GLY	TYR	GLY	ASN	THR	THR	VAL	ASN	SER	VAL	PRO	ASP	PHE	ASP	ASP	ALA	ASN	LYS	T
AACGAGGTATACCGTAAACACAACAGTAAATTCTGTAACCTGATTTGACGATGCAAAATAA																				82
1330																				1370
HR	LEU	ALA	ILE	THR	PHE	VAL	ASP	ALA	GLY	ARG	ARG	LEU	THR	VAL	ARG	GLN	LEU	ARG	P	
CATTAGCGATAACCTTTGTTGATGCTGGACAGCTTAACTGTTAACCTGCTTACCTGCCT																				
1390																				1440
HE	GLU	GLY	ASN	THR	VAL	SER	ALA	ASP	SER	THR	LEU	ARG	GLN	MET	ARG	GLN	GLN	GLU	G	
TTGAAAGGAAAATACCGTTRCTGCTGATAGTACTTACCGTACGGAATGCGCACAAAGAAG																				
1450																				1500
1460																				
1470																				
1480																				
1490																				

## FIG.1E.(CONTINUED)

LY	THR	TRP	TYR	ASN	SER	GIN	LEU	VAL	GLU	LEU	GLY	LYS	ILE	ARG	LEU	ASP	ARG	THR	GLY	P
G	A	A	C	T	T	G	G	T	A	T	A	A	T	C	G	C	T	A	G	T
1510	1520	1530	1540	1550	1560															
HE	PHE	GLU	THR	VAL	GLU	ASN	ARG	ILE	ASP	PRO	ILE	ASN	GLY	SER	ASN	ASP	GLU	VAL	ASP	V
T	C	T	T	C	G	A	A	C	A	G	T	T	G	A	T	T	G	G	A	T
1570	1580	1590	1600	1610	1620															
AL	VAL	TYR	LYS	VAL	LYS	GLU	ARG	ASN	THR	GLY	SER	ILE	ASN	PHE	GLY	IIE	GLY	TYR	GLY	T
T	C	G	T	A	T	A	A	G	T	C	A	C	G	G	T	A	C	T	T	G
1630	1640	1650	1660	1670	1680															
HR	GLU	SER	GLY	ILE	SER	TYR	GLN	THR	SER	ILE	LYS	GLN	ASP	ASN	PHE	LEU	GLY	THR	GLY	A
C	A	G	A	G	T	G	G	T	A	T	C	A	A	G	T	A	T	T	G	G
1690	1700	1710	1720	1730	1740															
IA	ALA	VAL	SER	ILE	ALA	GLY	THR	LYS	ASN	ASP	TYR	GLY	THR	SER	VAL	ASN	LEU	GLY	TYR	T
C	G	G	C	A	G	T	A	G	T	A	T	G	G	T	A	T	T	G	G	T
1750	1760	1770	1780	1790	1800															
HR	GLU	PRO	TYR	PHE	THR	LYS	ASP	GLY	VAL	SER	LEU	GLY	GLY	ASN	ILE	PHE	PHE	GLU	ASN	T
C	C	G	A	A	C	C	T	A	T	T	A	G	T	G	T	C	T	T	G	A
1810	1820	1830	1840	1850	1860															

SUBSTITUTE SHEET

39/82

FIG. 1E. (CONTINUED)

## **SUBSTITUTE SHEET**

## FIG.1E.(CONTINUED)

SN GLY PHE GLY ASN LYS ARG LEU PRO PHE TYR GLN THR TYR THR ALA GLY ILE GLY S  
 A T G G T T T G G A A C A A G C G T T A C C G T T C T A C T C A A C T T A C A G C G G T G G C A T T G G T T 2340  
 2290 2300 2310 2320 2330  
 ER LEU ARG GLY PHE ALA TYR GLY SER ILE GLY PRO ASN ALA ILE TYR ALA GLU HIS GLY A  
 C A T T A C C G C G G T T G C T T A T G G T A G T A T G C A A T T G C A A T T G C C G A A C A T G G T A 2400  
 2350 2360 2370 2380 2390  
 SN GLY THR PHE ASN LYS ILE SER ASP VAL ILE GLY GLY ASN ALA ILE THR THR ALA S  
 A T G G T A C T T T A A T A A G A T A A G T T C T G A T G T G A T T G G T A A T G C A A T C A C A A C T G C G A 41/82  
 2410 2420 2430 2440 2450  
 ER ALA GLU LEU ILE VAL PRO THR PRO PHE VAL SER ASP IYS SER GIN ASN THR VAL ARG T  
 G T G C A G A A C T T A T T G T A C C A A C T C C A T T G T G A G T G A T A A A A G C C C A A A T A C A G T C C G A A 2460  
 2470 2480 2490 2500 2510  
 HR SER LEU PHE VAL ASP ALA ALA SER VAL TRP ASN THR LYS TRP LYS SER ASP LYS ASN G  
 C C T C C C T A T T G T G A T G C G G C A A G T G T T G G A A T A C T A A A T G G C A A A T C A G A T A A A A T G 2520  
 2530 2540 2550 2560 2570  
 LY LEU GLU SER LYS VAL LEU LYS ASP LEU PRO ASP TYR GLY LYS SER SER ARG ILE ARG A  
 G A T T A G A G C A A G G C T T G A A A G A C T T A C C T G A T T A T G G C A A A T C A A G C C G T A T T C G C G 2640  
 2590 2600 2610 2620 2630

## FIG.1E.(CONTINUED)

LA SER THR GLY VAL GLY PHE GIN TRP GIN SER PRO ILE GLY PRO LEU VAL PHE SER TYR A  
 CCTCTACAGGTGGATTCGAAATGGCAATCTCCTATGGACCATTGGTATTCTTATG  
 2650 2660 2670 2680 2690 2700 2700  
 LA LYS PRO ILE LYS LYS TYR GLU ASN ASP VAL GLU GIN PHE GLN PHE SER ILE GLY G  
 CTAAACCAATTAAATAATGAAATGATGATGTCGAACAGTTCCAAATTAGTATTGGGG  
 2710 2720 2730 2740 2750 2760 2760  
 LY SER PHE \*\*\* \* \* \*  
 GCTCTTCTAAATAATTGAAACTTTCGTCATCAGAACCTCAAGAAAAACGACGTTCTGCC  
 2770 2780 2790 2800 2810 2820 2820  
 42/82  
 TAAATTGAAATTGGCGAGAAAATATAACCCATTAATTAGGATAATTCAAAAT  
 2830 2840 2850 2860 2870 2880 2880  
 GAAAAACATCGCAAAAGTAACCGCACCTTGCTTAGGTTTGCACTTGCTCAGGCTATGC  
 2890 2900 2910 2920 2930 2940 2940  
 TTCCGGCTGAAGAAAAATTGCTTCAATTATGCAGGTATATTCAA  
 2950 2960 2970 2980 2980 2990 2990

43/82

FIG. 1 F.

1. cad15  
3. minnad15  
2. eagand15  
4. pakd15  
5. sb33d15

cad15 1

1  
minnad15

eagannd15 1

pakd15

Sb33d15

consensus

1  
cad15

aaaaaggcattgaaaaacaggacagctttccctttaaccttggaaatattagggaaaatt

ACAGGACAGCTTCCCTTTAACCTTGAAATATTAGGGAAATT  
| | | | | | | | | | | | | | | | | | | | | | | |  
GGCATTGAAAAACAGGACAACTTCCCTTTAACCTTGAAATATTAGGGAAATT  
| | | | | | | | | | | | | | | | | | | | | | | |  
GGCATTGAAAAACAGGACAGCTTCCCTTTAACCTTGAAATATTAGGGAAATT

FIG. 1F. (CONTINUED)

minnad15	6	CTTactGGCGATTGTCAATTAAATAATTAAAGTGGGCCAATTCTATTGCAAAGGTGCTG	
eagand15	46	CTTcCCTGGCGATTGTCAATTAAATAATTAAAGTGGGCCAATTCTATTGCAAAGGTGCTG	
pakd15	62	CTTACTGGCGATTGTCAATTAAATAATTAAAGTGGGCCAATTCTATTGCAAAGGTGCTG	
sb333d15	58	CTTACTGGCGATTGTCAATTAAATAATTAAAGTGGGCCAATTCTATTGCAAAGGTGCTG	
consensus			
cad15	1	cttactggcgatttgtcattaataatctaaggccaaattttctatttgcaaaaagggtctg	
minnad15	67	GCaCATCAGCAAATATTGGATTTGGTGTATTTTAAGTTTATGGCACTGATTAGTGTAAA	
eagand15	107	GCCCATCAGCAAATATTGGATTTGGTGTATTTTAAGTTTATGGCACTGATTAGTGTAAA	
pakd15	123	GtGCATCAGCAAATATTGGATTTGGTGTATTTTAAGTTTATGGCATTGATTAGTGTAAA	
sb333d15	119	GCGCATTAGCAAATATTGGATTTGGTGTATTTTAAGTTTATGGCATTGATTAGTGTAAA	
consensus			

FIG. 1F. (CONTINUED)

	cad15	1	gATTAC	
minnad15	128	TTTAGGGATTATGAATTATTCCATTACCACTTATTAGATGGGGTCATTAGTTTTTA		
eagand15	168	TTTAGGGATTATGAATTATTCCATTACCACTTATTAGATGGGGTCATTAGTTTTTA		
pakd15	184	TTTAGGGATTATGAATTATTCCATTACCACTTATTAGATGGGGTCATTAGTTTTTA		
sb33d15	180	TTTAGGGATTATGAATTATTCCATTACCACTTATTAGATGGGGTCATTAGTTTTTA		
consensus				
	cad15	7		
minnad15	189	ACAATGGAAGCTGTTAAGAAAACCTGTTCTGAGGGGTGCAAAGCATCTGTTATCGAA		
eagand15	229	ACAATGGAAGCTGTTAAGAAAACCTGTTCTGAGGGGTGCAAAGCATCTGTTATCGAA		
pakd15	245	ACAATGGAAGCTGTTAAGAAAACCTGTTCTGAGGGGTGCAAAGCATCTGTTATCGAA		
sb33d15	241	ACAATGGAAGCTGTTAAGAAAACCTGTTCTGAGGGGTGCAAAGCATCTGTTATCGAA		

## FIG. 1F. (CONTINUED)

consensus

acaatggaaaggtaaaaggaaaaaccctgttctgagggggtgcaaaagcatctgttatcgaa

cad15	7	gccaAGCTTAACGGTGTTCGATTATTGATTTTACGTCT	
minnad15	250	TTGGCGCAGCACTGTTAACGGCTTAAAGCTTAAATGATTATTGCATTACGTCT	
eagand15	290	TTGGCGCAGCACTGTTAACGGCTTAAAGCTTAAATGATTATTGCATTACGTCT	
pakd15	306	TTGGCGCAGCACTGTTAACGGCTTAAAGCTTAAATGATTATTGCATTACGTCT	
sb33d15	302	TTGGCGCAGCACTGTTAACGGCTTAAAGCTTAAATGATTATTGCATTACGTCT	
consensus		ttggccgcaggcactgttattAAAGCTTAACGGTGTTCGATTATGATTACGTCT	46/82
cad15	52	ATAAATTATAAGGATAACAATCGATGAAAAAAACTCTAAATCGCAAGTTATTACGGTAC	
minnad15	311	ATAAATTATAAGGATAACAATCGATGAAAAAAACTCTAAATCGCAAGTTATTACGGTAC	
eagand15	351	ATAAATTATAAGGATAACAATCGATGAAAAAAACTCTAAATCGCAAGTTATTACGGTAC	
pakd15	367	ATAAATTATAAGGATAACAATCGATGAAAAAAACTCTAAATCGCAAGTTATTACGGTAC	

## FIG.1F.(CONTINUED)

sb33d15	363	ATAAATTATAGGATAACAATCGATGAAAAAAACTTCTAATCGCAAGTTATTTCGGT <sup>a</sup> C	
consensus			
cad15	113	GACAAACGACTGTGTTGCCGCACCTTTGTGGCAAAAGATAATTCGTGTGGATGGTGTCAA	
minnad15	372	GACAAACGACTGTGTTGCCGCACCTTTGTGGCAAAAGATAATTCGTGTGGATGGTGTCAA	
eagand15	412	GACAAACGACTGTGTTGCCGCACCTTTGTGGCAAAAGATAATTCGTGTGGATGGTGTCAA	
pakd15	428	GACAAACGACTGTGTTGCCGCACCTTTGTGGCAAAAGATAATTCGTGTGGATGGTGTCAA	<sup>47/82</sup>
sb33d15	424	GACAAACGACTGTGTTGCCGCACCTTTGTGGCAAAAGATAATTCGTGTGGATGGTGTCAA	
consensus			
cad15	174	GGTGACTTAGAACAAATCCGAGCAAGTTACCTGTCGTGCCGGTCAGCGTGTGACTG	
minnad15	433	GGTGACTTAGAACAAATCCGAGCAAGTTACCTGTCGTGCCGGTCAGCGTGTGACTG	
eagand15	473	GGTGACTTAGAACAAATCCGAGCAAGTTACCTGTCGTGCCGGTCAGCGTGTGACTG	

## FIG. 1F. (CONTINUED)

pakd15	489	GGTGA <del>CT</del> TAGAACAAATCCGAGCAAGTTACCTGTTCGTGCTGGTCAGCGTGTGACTG	
sb33d15	485	GGTGA <del>CT</del> TAGAACAAATCCGAGCAAGTTACCTGTTACCTGTTCGTGCCGGTCAGCGTGTGACTG	
consensus			
GGTGA <del>CT</del> TAGAACAAATCCGAGCAAGTTACCTGTTCGTGCCGGTCAGCGTGTGACTG			
cad15	235	ACAATGATGGCTAATAATTGCCGCTCTTATTCGTAAGTGGTCGATTGATGATGTGAA	
minnad15	494	ACAATGATGGCTAATAATTGCCGCTCTTATTGTAAGTGGTCGATTGATGATGTGAA	
eagand15	534	ACAATGATGGCTAATAATTGCCGCTCTTATTGTAAGTGGTCGATTGATGATGTGAA	<sup>48/82</sup>
pakd15	550	ACAATGATGGCTAATAATTGCCGCTCTTATTGTAAGTGGTCGATTGATGATGTGAA	
sb33d15	546	ACAATGATGGCTAATAATTGCCGCTCTTATTGTAAGTGGTCGATTGATGATGTGAA	
consensus			
ACAATGATGGCTAATAATTGCCGCTCTTATTGTAAGTGGTCGATTGATGATGTGAA			
cad15	296	AGGGCATCAAAGAAGGGGATGGCTTGTAGCGTTGCTAAATCGATCATTAGAT	
minnad15	555	AGGGCATCAAAGAAGGGGATGGCTTGTAGCGTTGCTAAATCGATCATTAGAT	
consensus			
AGGGCATCAAAGAAGGGGATGGCTTGTAGCGTTGCTAAATCGATCATTAGAT			

FIG. 1F. (CONTINUED)

				49/82
eagand15	595	ACGGCATCAAGAAGGGATGTCCTGTGGCTAAATCGATCATTAGATGCTA		
pakd15	611	ACGGCATCAAGAAGGGATGTCCTGTGGCTAAATCGATCATTAGATGCTA		
sb33d15	607	ACGGCATCAAGAAGGGATGTCCTGTGGCTAAATCGATCATTAGATGCTA		
consensus		ACGGCATCAAGAAGGGATGTCCTGTGGCTAAATCGATCATTAGATGCTA		
cad15	357	GTTAAATCAAAGGTAAACTCTGTATTCCCAC'TGAAGCACTAAACAAACTTAGATGCTA		
minnad15	616	GTTAAATCAAAGGTAACTCTGTATTCCCAC'TGAAGCACTAAACAAACTTAGATGCTA		
eagand15	656	GTTAAATCAAAGGTAACTCTGTATTCCCAC'TGAAGCACTAAACAAACTTAGATGCTA		
pakd15	672	GTTAAATCAAAGGTAACTCTGTATTCCCAC'TGAAGCACTAAACAAACTTAGATGCTA		
sb33d15	668	GTTAAATCAAAGGTAACTCTATTCCac'CTGAAGCACTAAACAAACTTAGATGCTA		
consensus		GTTAAATCAAAGGTAACTCTGTATTCCCac'CTGAAGCACTAAACAAACTTAGATGCTA		
cad15	418	ACGGGTTAAAGTTGGCGATGTTAATTGAGAAAATTAAATGAATTGCCAAAAGTGT		

FIG. 1F. (CONTINUED)

minnad15	677	ACGGGT'TTAAAGTTGGCGATGTTTAATTGAAATTGAGAAAATTGAAATTGAGATGTTAATTGCCAAAGTGT
eagand15	717	ACGGGT'TTAAAGTTGGCGATGTTTAATTGAGAAAATTGAAATTGAGATGTTAATTGCCAAAGTGT
pakd15	733	ACGGGT'TTAAAGTTGGCGATGTTAATTGAAATTGAGAAAATTGAAATTGAGATGTTAATTGCCAAAGTGT
sb333d15	729	ACGGGT'TTAAAGTTGGCGATATTAAATTGAAATTGAGAAAATTGAAATTGAGATTTGCCAAAGTGT
consensus		
cad15	479	AAAAGAGCACTATGCAAGTGTAGGTGGCTATAACGCAAACAGTTGAACCTATTGTCAATAACG
minnad15	738	AAAAGAGCACTATGCAAGTGTAGGTGGCTATAACGCAAACAGTTGAACCTATTGTCAATAACG
eagand15	778	AAAAGAGCACTATGCAAGTGTAGGTGGCTATAACGCAAACAGTTGAACCTATTGTCAATAACG
pakd15	794	AAAAGAGCACTATGCAAGTGTAGGTGGCTATAACGCAAACGGTTGAACCTATTGTCAATAACG
sb333d15	790	AAAAGAGCACTATGCAAGTGTAGGTGGCTATAACGCAAACGGTTGAACCTATTGTCAATAACG
consensus		

FIG. 1F. (CONTINUED)

			540	CTACCAATAATCGGCTGAAATTAAATTCAAATGAAGATGATAAGCAAATTTGG	51/82
minnad15	799			CTACCAATAATCGGCTGAAATTAAATTCAAATGAAGATGATAAGCAAATTTGG	
eagand15	839			CTACCAATAATCGGCTGAAATTAAATTCAAATGAAGATGATAAGCAAATTTGG	
pakd15	855			CTgCCAAATAATCGTGCTGAAATTAAATTCAAATGAAGATGATAAGCAAATTTGG	
sb33d15	851			CTACCAATAATCGGCTGAAATTAAATTCAAATGAAGATGATAAGCAAATTTGG	
consensus				CTACCAATAATCGGCTGAAATTAAATTCAAATGAAGATGATAAGCAAATTTGG	
cad15	601			CATCATTAACTTCAAGGGAACGAAACTGTAGCTGAGTACATTACAAGAACAAATGGA	
minnad15	860			CATCATTAACTTCAAGGGAACGAAACTGTAGCTGAGTACATTACAAGAACAAATGGA	
eagand15	900			CATCATTAACTTCAAGGGAACGAAACTGTAGCTGAGTACATTACAAGAACAAATGGA	
pakd15	916			CATCATTAACTTCAAGGGAACGAAACTGTAGCTGAGTACATTACAAGAACAAATGGA	
sb33d15	912			CATCATTAACTTCAAGGGAACGAAACTGTAGCTGAGTACATTACAAGAACAAATGGA	
consensus				CATCATTAACTTCAAGGGAACGAAACTGTAGCTGAGTACATTACAAGAACAAATGGA	

FIG. 1F. (CONTINUED)

cad15	662	ATTACAAACCTGATTCTTGGTGGAAATTATGGGAAATAAATTGAAGGTGCCCAATTTCGAG
minnad15	921	ATTACAAACCTGATTCTTGGTGGAAATTATGGGAAATAAATTGAAGGTGCCCAATTTCGAG
eagand15	961	ATTACAAACCTGATTCTTGGTGGAAATTATGGGAAATAAATTGAAGGTGCCCAATTTCGAG
pakd15	977	ATTACAAACCTGATTCTTGGTGGAAATTATGGGAAATAAATTGAAGGTGCCCAATTTCGAG
sb33d15	973	ATTACAAACCTGATTCTTGGTGGAAATTATGGGAAATAAATTGAAGGTGCCCAATTTCGAG

62/82

			consensus
cad15	723	AAAGATTGCACTCGTGAATTATTAAATAATGGCTATGCCAAGGCACAAATTAA	
minnad15	982	AAAGATTGCACTCGTGAATTTCGTGATTATTAAATAATGGCTATGCCAAGGCACAAATTAA	
eagand15	1022	AAAGATTGCACTCGTGAATTTCGTGATTATTAAATAATGGCTATGCCAAGGCACAAATTAA	
pakd15	1038	AAAGATCTGAGGCCATTCTCGTGATTATTAAATAATGGCTATGCCAAGGCACAAATTCA	
sb33d15	1034	AAAGATTGCACTCGTGAATTTCGTGATTATTAAATAATGGCTATGCCAAGGCACAAATTCA	
			AAAGATTGCACTCGTGAATTATTAAATAATGGCTATGCCAAGGCACAAATTAA

## FIG.1F.(CONTINUED)

53/82

cad15	784	CTAAAACGGATGTTCAAGCTAAATGATGAAAAAACAAAAGTTAATGTAACCATTGATGTAAA	
minnad15	1043	CTAAAACGGATGTTCAAGCTAAATGATGAAAAAACAAAAGTTAATGTAACCATTGATGTAAA	
eagand15	1083	CTAAAACGGATGTTCAAGCTAAATGATGAAAAAACAAAAGTTAATGTAACCATTGATGTAAA	
pakd15	1099	CTAAAACGGATGTTCAAGCTAAATGATGAAAAAACAAAAGTTAATGTAACCATTGATGTAAA	
sb33d15	1095	CTAAAACGGATGTTCAAGCTAAATGATGAAAAAACAAAAGTTAATGTAACCATTGATGTAAA	
consensus			
cad15	845	TGAAGGTTTACAGTATGACCTTCGTAGTGCACGCATTATAGGTAATCTGGAGGTATGTCT	
minnad15	1104	TGAAGGTTTACAGTATGACCTTCGTAGTGCACGCATTATAGGTAATCTGGAGGTATGTCT	
eagand15	1144	TGAAGGTTTACAGTATGACCTTCGTAGTGCACGCATTATAGGTAATCTGGAGGTATGTCT	
pakd15	1160	TGAAGGTTTACAGTATGACCTTCGTAGTGCACGCATTATAGGTAATCTGGAGGTATGTCT	
sb33d15	1156	TGAAGGTTTACAGTATGACCTTCGTAGTGCACGCATTATAGGTAATCTGGAGGTATGTCT	
consensus			

## FIG. 1F.(CONTINUED)

cad15	906	GCCGAGCTTGAACCTTACTTTCAGCATTACATTAAATGATACTTACATTAAATGATACTTTACGGCTTAGTGATA
minnad15	1165	GCCGAGCTTGAACCTTACTTTCAGCATTACATTAAATGATACTTACATTAAATGATACTTTACGGCTTAGTGATA
eagand15	1205	GCCGAGCTTGAACCTTACTTTCAGCATTACATTAAATGATACTTACATTAAATGATACTTTACGGCTTAGTGATA
pakd15	1221	GCCGAGCTTGAACCTTACTTTCAGCATTACATTAAATGATACTTACATTAAATGATACTTTACGGCTTAGTGATA
sb33d15	1217	GCCGAGCTTGAACCTTACTTTCAGCATTACATTAAATGATACTTACGGCTTAGTGATA
consensus		GGCGAGCTTGAACCTTACTTTCAGCATTACATTAAATGATACTTACGGCTTAGTGATA

54/82

cad15	967	TTGCAGATGTAGAAAATGCAATTAAAGCAAACCTGGAGAACGGGGTTACGGTAGGGCAAC
minnad15	1226	TTGCAGATGTAGAAAATGCAATTAAAGCAAACCTGGAGAACGGGGTTACGGTAGGGCAAC
eagand15	1266	TTGCAGATGTAGAAAATGCAATTAAAGCAAACCTGGAGAACGGGGTTACGGTAGGGCAAC
pakd15	1282	TTGCAGATGTAGAAAATGCAATTAAAGCAAACCTGGGAACGAGGTTACGGTAACACAAAC
sb33d15	1278	TTGCAGATGTAGAAAATGCAATTAAAGCAAACCTGGGAACGAGGTTACGGTAACACAAAC
consensus		TTGCAGATGTAGAAAATGCAATTAAAGCAAACCTGGAGAACGGGGTTACGGTAGGGCAAC

55/82

FIG. 1F. (CONTINUED)

cad15	1028	GGTAAATTCACTGATTGATGATAACCATAGCGATAACCCTTGTGTT
minnad15	1287	GGTAAATTCACTGATTGATGATAACCATAGCGATAACCCTTGTGTT
eagand15	1327	GGTAAATTCACTGATTGATGATAACCATAGCGATAACCCTTGTGTT
pakd15	1343	AGTAAATTCTGACTGATTGACGATGCAAATAACCATAGCGATAACCCTTGTGTT
sb33d15	1339	AGTAAATTCTGACTGATTGACGATGCAAATAACCATAGCGATAACCCTTGTGTT

			consensus
cad15	1089	GATGCTGGACGACGTAACTTGCACCAACTTCGCTTTGAAGGAAATAACCGTTCTGCTG	
minnad15	1348	GATGCTGGACGACGTAACTTGCACCAACTTCGCTTTGAAGGAAATAACCGTTCTGCTG	
eagand15	1388	GATGCTGGACGACGTAACTTGCACCAACTTCGCTTTGAAGGAAATAACCGTTCTGCTG	
pakd15	1404	GATGCTGGACGACGTAACTTGCACCAACTTCGCTTTGAAGGAAATAACCGTTCTGCTG	
sb33d15	1400	GATGCTGGACGACGTAACTTGCACCAACTTCGCTTTGAAGGAAATAACCGTTCTGCTG	
			consensus

FIG. 1F. (CONTINUED)

			consensus
cad15	1150	ATAGCACTTACGT CAGGAAATGGGCCAACAGAACGAAACTGGTATAATTCAAATTAGT	
minnad15	1409	ATAGCACTTACGT CAGGAAATGGGCCAACAGAACGAAACTGGTATAATTCAAATTAGT	
eagand15	1449	ATAGCACTTACGT CAGGAAATGGGCCAACAGAACGAAACTGGTATAATTCAAATTAGT	
pakd15	1465	ATAGTACTTACGT CAGGAAATGGGCCAACAGAACGAAACTGGTATAATTCAAATTAGT	
sb33d15	1461	ATAGTACTTACGT CAGGAAATGGGCCAACAGAACGAAACTGGTATAATTCAAATTAGT	
			ATAGCACTTACGT CAGGAAATGGGCCAACAGAACGAAACTGGTATAATTCAAATTAGT

consensus

					TGAGT'TAGGAAAAT'CGCTTAGATCGTACAGGT'TCTTCGAAACAGTCGAAACCGAATT
cad15	1211				TGAGT'TAGGAAAAT'CGCTTAGATCGTACAGGT'TCTTCGAAACAGTCGAAACCGAATT
minnad15	1470				TGAGT'TAGGAAAAT'CGCTTAGATCGTACAGGT'TCTTCGAAACAGTCGAAACCGAATT
eagand15	1510				TGAGT'TAGGAAAAT'CGCTTAGATCGTACAGGT'TCTTCGAAACAGTCGAAACCGAATT
pakd15	1526				TGAGT'TAGGAAAAT'CGCTTAGATCGTACAGGT'TCTTCGAAACAGTCGAAACCGAATT
sb33d15	1522				TGAGT'TAGGAAAAT'CGCTTAGATCGTACAGGT'TCTTCGAAACAGTCGAAACCGAATT
					TGAGT'TAGGAAAAT'CGCTTAGATCGTACAGGT'TCTTCGAAACAGTCGAAACCGAATT
					consensus

FIG. 1F. (CONTINUED)

cad15	1272	GATCCTATCAATGGTAGTAATGATGAAGTGGATGTCGTATAAAGTCAAAGAACGTTAAC
minnad15	1531	GATCCTATCAATGGTAGTAATGATGAAGTGGATGTCGTATAAAGTCAAAGAACGTTAAC
eagand15	1571	GATCCTATCAATGGTAGTAATGATGAAGTGGATGTCGTATAAAGTCAAAGAACGTTAAC
pakd15	1587	GATCCTATCAATGGTAGCAATTGATGAAGTGGATGTCGTATAAAGTCAAAGAACGTTAAC
sb3 3d15	1583	GATCCTATCAATGGTAGCAATTGATGAAGTGGATGTCGTATAAAGTCAAAGAACGTTAAC

consensus

57/82

cad15	1333	CGGGTAGTATACTCAACTTTGGTATTGGTACGGTACAGAGAGTGGTATTAGTTATCAAGCAAG
minnad15	1592	CGGGTAGTATACTCAACTTTGGTATTGGTACGGTACAGAGAGTGGTATTAGTTATCAAGCAAG
eagand15	1632	CGGGTAGTATACTCAACTTTGGTATTGGTACGGTACAGAGAGTGGTATTAGTTATCAAGCAAG
pakd15	1648	CGGGTAGTATACTCAACTTTGGTATTGGTACGGTACAGAGAGTGGTATCAGTTATCAAACAAG
sb33d15	1644	CGGGTAGTATACTCAACTTTGGTATTGGTACGGTACAGAGAGTGGTATTAGTTATCAAGCAAG

CGGGTAGTATCACTTTGGTATTGGTTACGGTACAGAGGTATTAGTTATCAAGCAAG  
consensus

## FIG.1F.(CONTINUED)

8/82

cad15	1394	TGTAAACAGATAATTCTTGGAACAGGGGGCAGTAAGTATAGCTGGTACGAAAT	
minnad15	1653	TGTTAAACAGATAATTCTTGGAACAGGGGGCAGTAAGTATAGCTGGTACGAAAT	
eagand15	1693	TGTTAAACAGATAATTCTTGGAACAGGGGGCAGTAAGTATAGCTGGTACGAAAT	
pakd15	1709	TATTAACAGATAATTCTTGGAACAGGGGGCAGTAAGTATAGCTGGTACGAAAT	
sb33d15	1705	TgtcAAACAGATAATTCTTGGAACAGGGGGCAGTAAGTATAGCTGGTACGAAAT	
consensus			
cad15	1455	GATTATGGTACGAGTGTCAATTGGTTATAACCGAGGCCCTATTACTAAAGATGGTGTAA	
minnad15	1714	GATTATGGTACGAGTGTCAATTGGTTATAACCGAGGCCCTATTACTAAAGATGGTGTAA	
eagand15	1754	GATTATGGTACGAGTGTCAATTGGTTATAACCGAGGCCCTATTACTAAAGATGGTGTAA	
pakd15	1770	GATTATGGTACGAGTGTCAATTGGTTATAACCGAacCCCTATTACTAAAGATGGTGTAA	
sb33d15	1766	GATTATGGTACGAGTGTCAATTGGTTATAACCGAGGCCCTATTACTAAAGATGGTGTAA	
consensus			

## FIG.1F.(CONTINUED)

cad15	1516	GTCTTGGAAATGTTCTTGAAGACTACGATAACTCTAAAAGTGATAACATCCTCTAA
minnad15	1775	GTCTTGGAAATGTTCTTGAAGACTACGATAACTCTAAAAGTGATAACATCCTCTAA
eagand15	1815	GTCTTGGAAATGTTCTTGAAGACTACGATAACTCTAAAAGTGATAACATCCTCTAA
pakd15	1831	GTCTTGGAAATTTCTTGTAAACTACGATAACTCTAAAAGTGATAACATCCTCTAA
sb33d15	1827	GTCTTGGAAATGTTCTTGTAAACTACGATAACTCTAAAAGTGATAACATCCTCTAA
		consensus
GTCTTGGAAATGTTCTTGTAAACTACGATAACTCTAAAAGTGATAACATCCTCTAA		

59/82

consensus

cad15	1577	CTATAAGCGTACGACTTACGGAAAGTAATGTTACTTTAGGTTCCTGTAATGAAAATAAAC
minnad15	1836	CTATAAGCGTACGACTTACGGAAAGTAATGTTACTTTAGGTTCCTGTAATGAAAATAAAC
eagand15	1876	CTATAAGCGTACGACTTACGGAAAGTAATGTTACTTTAGGTTCCTGTAATGAAAATAAAC
pakd15	1892	CTATAAGCGTACGACTTACGGAAAGTAATGTTACTTTAGGTTCCTGTAATGAAAATAAAC
sb33d15	1888	CTATAAGCGTACGACTTACGGAAAGTAATGTTACTTTAGGTTCCTGTAATGAAAATAAAC
		consensus
CTATAAGCGTACGACTTACGGAAAGTAATGTTACTTTAGGTTCCTGTAATGAAAATAAAC		

## FIG.1F.(CONTINUED)

60/82

cad15	1638	TCCTATTAGGATTAACCTATAAAATTAGTAACTTGGCTCTAGAATA	
minnad15	1897	TCCTATTAGGATTAACCTATAAAATTAGTAACTTGGCTCTAGAATA	
eagand15	1937	TCCTATTAGGATTAACCTATAAAATTAGTAACTTGGCTCTAGAATA	
pakd15	1953	TCCTATTAGGATTAACCTATAAAATTAGTAACTTGGCTCTAGAATA	
sb33d15	1949	TCCTATTAGGATTAACCTATAAAATTAGTAACTTGGCTCTAGAATA	
		consensus	TCCTATTAGGATTAACCTATAAAATTAGTAACTTGGCTCTAGAATA
cad15	1699	ACCGTAATTATTCAAATCAAATGAAATTAAAGGTAATGGCATTAAACAAATGACTT	
minnad15	1958	ACCGTAATTATTCAAATCAAATGAAATTAAAGGTAATGGCATTAAACAAATGACTT	
eagand15	1998	ACCGTAATTATTCAAATCAAATGAAATTAAAGGTAATGGCATTAAACAAATGACTT	
pakd15	2014	ACCGTAATTATTCAAATCAAATGAAATTAAAGGTAATGGCATTAAACAAATGACTT	
sb33d15	2010	ACCGTAATTATTCAAATCAAATGAAATTAAAGGTAATGGCATTAAACAAATGACTT	
		consensus	ACCGTAATTATTCAAATCAAATGAAATTAAAGGTAATGGCATTAAACAAATGACTT

FIG. 1F. (CONTINUED)

cad15	1760	TGATTTTCTTGGAACTATAACAGCCTTAATAGAGGCATTTCACAACTAAAGGG			
minnad15	2019	TGATTTTCTTGGTTGGAACTATAACAGCCTTAATAGAGGCATTTCACAACTAAAGGG			
eagand15	2059	TGATTTTCTTGGTTGGAACTATAACAGCCTTAATAGAGGCATTTCACAACTAAAGGG			
pakd15	2075	TGATTTTCTTGGTTGGAACTATAACAGCCTTAATAGAGGCATTTCACAACTAAAGGG			
sb33d15	2071	TGATTTTCTTGGTTGGAACTATAACAGCCTTAATAGAGGCATTTCACAACTAAAGGG			
			consensus		
			TGATTTTCTTGGTTGGAACTATAACAGCCTTAATAGAGGCATTTCACAACTAAAGGG		
				61/82	
cad15	1821	GTTAAAGCAAGTCTTGGGACGAGGTACTATCCAGGTTCTGATAACAAATACTACAAAC			
minnad15	2080	GTTAAAGCAAGTCTTGGGACGAGGTACTATCCAGGTTCTGATAACAAATACTACAAAC			
eagand15	2120	GTTAAAGCAAGTCTTGGGACGAGGTACTATCCAGGTTCTGATAACAAATACTACAAAC			
pakd15	2136	GTTAAAGCAAGTCTTGGGACGAGGTACTATCCAGGTTCTGATAACAAATACTACAAAC			
sb33d15	2132	GTTAAAGCAAGTCTTGGGACGAGGTACAATCCAGGTTCTGATAACAAATACTACAAAC			
			consensus		
			GTTAAAGCAAGTCTTGGGACGAGGTACTATCCAGGTTCTGATAACAAATACTACAAAC		

## FIG.1F.(CONTINUED)

						62/82
cad15	1882	TAAGTGCAGATGTACAGGGTTCTACCCATTAGACAGAGATCACCTCTGGGTTGTATCTGC				
minnad15	2141	TAAGTGCAGATGTACAGGGTTCTACCCATTAGACAGAGATCACCTCTGGGTTGTATCTGC				
eagand15	2181	TAAGTGCAGATGTACAGGGTTCTACCCATTAGACAGAGATCACCTCTGGGTTGTATCTGC				
pakd15	2197	TAAGTGCAGATGTACAGGGTTCTACCCATTAGACAGAGATCACCGCTGGGTTGTATCTGC				
sb33d15	2193	TAAGTGCAGATGTACAGGGTTCTACCCATTAGACAGAGATCACCTCTGGGTTGTATCTGC				
				consensus	TAAGTGCAGATGTACAGGGTTCTACCCATTAGACAGAGATCACCTCTGGGTTGTATCTGC	
cad15	1943	AAAAGCATTCTGCAGGATATGCAAATGGTTTGAAACAAGCGTTACCGTTCTATCAAAC				
minnad15	2202	AAAAGCATTCTGCAGGATATGCAAATGGTTTGAAACAAGCGTTACCGTTCTATCAAAC				
eagand15	2242	AAAAGCATTCTGCAGGATATGCAAATGGTTTGAAACAAGCGTTACCGTTCTATCAAAC				
pakd15	2258	AAAAGCATTCTGCAGGATATGCAAATGGTTTGAAACAAGCGTTACCGTTCTATCAAAC				
sb33d15	2254	AAAAGCATTCTGCAGGATATGCAAATGGTTTGAAACAAGCGTTACCGTTCTATCAAAC				

FIG. 1F. (CONTINUED)

consensus

AAAAGGCATCTGCAGGATATGCCAAATGGTTTGAAACAGCGTACCAACT

				6/82
cad15	2004	TATA CAG CGGGT GGCAT CGGTT CATTAC GTGGTT GCTT ATGGTAGT ATTGGACCTAACG		
minnad15	2263	TATA CAG CGGGT GGCAT CGGTT CATTAC GTGGTT GCTT ATGGTAGT ATTGGACCTAACG		
eagand15	2303	TATA CAG CGGGT GGCAT CGGTT CATTAC GTGGTT GCTT ATGGTAGT ATTGGACCTAACG		
pakd15	2319	TATA CAG CGGGT GGCAT CGGTT CATTAC GTGGTT GCTT ATGGTAGT ATTGGACCTAACG		
sb3d15	2315	TATA CAG CGGGT GGCAT CGGTT CATTAC GTGGTT GCTT ATGGTAGC ATTGGACCTAACG		
onsensus		TATA CAG CGGGT GGCAT CGGTT CATTAC GTGGTT GCTT ATGGTAGT ATTGGACCTAACG		

cad15	2065	CAATTATGCCGAATATGGTAATGGTAGTGGTACTGGTACTTTAAGAAAGATAAAGT'TCTGA
minnad15	2324	CAATTATGCCGAATATGGTAATGGTAGTGGTACTGGTACTTTAAGAAAGATAAAGT'TCTGA
eagand15	2364	CAATTATGCCGAATATGGTAATGGTAGTGGTACTGGTACTTTAAGAAAGATAAAGT'TCTGA
pakd15	2380	CAATTATGCCGAACATGGTAATGGTA

FIG. 1F. (CONTINUED)

consensus	CAATTATggTAATggtagtgtactggtaactTTAAGAAAGATAAAGTCTGA
sb33d15	2376 CAATTATcaAggtCAaaaATAAT
	aaATTAAAGATAAAGTCTGA

			TGTGATTGGTGGTAATGCCAATCGCTACAGCTAGGCAGAGTTAAATTGTGCCAACTCCATT
cad15	2126		TGTGATTGGTGGTAATGCCAATCGCTACAGCTAGGCAGAGTTAAATTGTGCCAACTCCATT
minnadi15	2385		TGTGATTGGTGGTAATGCCAATCGCTACAGCTAGGCAGAGTTAAATTGTGCCAACTCCATT
eagand15	2425		TGTGATTGGTGGTAATGCCAATCGCTACAGCTAGGCAGAGTTAAATTGTGCCAACTCCATT
pakd15	2429		TGTGATTGGTGGTAATGCCAATCGCTACAGCTAGGCAGAGTTAAATTGTGCCAACTCCATT
sb33d15	2422		TGTGATTGGTGGTAATGCCAATCGCTACAGCTAGGCAGAGTTAAATTGTGCCAACTCCATT
consensus			TGTGATTGGTGGTAATGCCAATCGCTACAGCTAGGCAGAGTTAAATTGTGCCAACTCCATT

cad15	2187	GTGAGCGATAAGGCCAAAATACGGTCCGAACCTCCTTATTGTTGATGCGGCCAAGTGT
minnad15	2446	GTGAGCGATAAGGCCAAAATACGGTCCGAACCTCCTTATTGTTGATGCGGCCAAGTGT
eagand15	2486	GTGAGCGATAAGGCCAAAATACGGTCCGAACCTCCTTATTGTTGATGCGGCCAAGTGT

## FIG.1F.(CONTINUED)

pakd15	2490	GTGAGTGTATAAAAGCCAAAATACAGTCCGAACCTCCTATTGTTGATGCCGGCAAGTGT		
sb33d15	2483	GTGAGTGTATAAGAGtCAAATACAGTCCGAACCTCCTATTGTTGATGCCGGCAAGTGT		
		<b>consensus</b>	GTGAGCGATAAAGAGCCAAATAAC9GTCCGAACCTCCTATTGTTGATGCCGGCAAGTGT	
cad15	2248	GGAAATACTAAATGGAATTCAGATAAAAATGGATTAGAGAGCGATGTATTAAAAAGATTGCC		
minnad15	2507	GGAAATACTAAATGGAATTCAGATAAAAATGGATTAGAGAGCGATGTATTAAAAAGATTGCC	65	/82
eagand15	2547	GGAAATACTAAATGGAATTCAGATAAAAATGGATTAGAGAGCGATGTATTAAAAAGATTGCC		
pakd15	2551	GGAAATACTAAATGGAATTCAGATAAAAATGGATTAGAGAGCGAAGGTCTTGAAAGACTTACC		
sb33d15	2544	GGAAATACTAAATGGAATTCAGATAAAAATGGATTAGAGAGCGAATGTCTTGAAAGACTTACC		
		<b>consensus</b>	GGAAATACTAAATGGAATTCAGATAAAAATGGATTAGAGAGCGATGTATTAAAAAGATTGCC	
cad15	2309	TGATTATGGCAAATCAAGCCGTATTTCGGCGCTCTACAGGTGTCGGATTCCAATGGCAATCT		
minnad15	2568	TGATTATGGCAAATCAAGCCGTATTTCGGCGCTCTACAGGTGTCGGATTCCAATGGCAATCT		

FIG.1F.(CONTINUED)

eagand15	2608	TGATTATGGCAAATCAAGGCCGTATT CGGCCTACAGGTGTCGGATTCCAATGGCAATCT
pakd15	2612	TGATTATGGCAAATCAAGGCCGTATT CGGCCTACAGGTGTCGGATTCCAATGGCAATCT
sb33d15	2605	CGATTATGGCAAATCAAGGCCGTACT CGGCCTACAGGTGTCGGATTCCAATGGCAATCT
consensus		tgattatggcaaatacgccgtatt cggctacagggtgtcggatccaatggcaatct

cad15 2431 TCGAACAGTTCCAATTTAGTATTGGAGGTTCTTAATAATTGAACCTTTCTTCATC

## FIG.1F.(CONTINUED)

minnad15	2690	TCGAACAGTCCAATTAGTATTGGAGGTTCTTAATAATTGAACTTTCTTCATC		
eagand15	2730	TCGAACAGTCCAATTAGTATTGGAGGTTCTTAATAATTGAACTTTCTTCATC		
pakd15	2734	TCGAACAGTCCAATTAGTATTGGGGCTCTTAATAAAATTGAACTTTCTGCATC		
sb33d15	2727	TCGAACAGTCCAATTAGTATTGGGGTTCTTAATAAAATTGAACTTTCTGCATC		
consensus				
67/82				
cad15	2492	AGAACTCTCAAACACGTTCTGCCTAATTAAATTGGGAGAGAAAATAAACCCATC		
minnad15	2751	AGAAACTCAAACACGTTCTGCCTAATTAAATTGGGAGAGAAAATAAACCCATC		
eagand15	2791	AGAAACTCTCAAACACGTTCTGCCTAATTAAATTGGGAGAGAAAATAAACCCATC		
pakd15	2795	AGAAACTCAAACACGAGACGTTCTGCCTAATTGAATTGGGAGAGAAAATAAACCCATC		
sb33d15	2788	AGAAACTCTCAAACGTTCTGCCTAATTAAATTGGGAGAGAAAATAAACCCATC		
consensus				

## FIG.1F.(CONTINUED)

cad15	2553	ATTTAATTAAAGGATATTCAAATGAAAACATCGCAAAAGTAACCGCACTTGCTTTAGG	68/82
minnad15	2812	ATTTAATTAAAGGATATTCAAATGAAAACATCGCAAAAGTAACCGCACTTGCTTTAGG	
eagand15	2852	ATTTAATTAAAGGATATTCAAATGAAAACATCGCAAAAGTAACCGCACTTGCTTTAGG	
pakd15	2856	ATTTAATTAAAGGATATTCAAATGAAAACATCGCAAAAGTAACCGCACTTGCTTTAGG	
sb33d15	2849	ATTTAATTAAAGGATATTCAAATGAAAACATCGCCAAAGTAACCGCACTTGCTTTAGG	
consensus		ATTTAATTAAAGGATATTCAAATGAAAACATCGCaAAAGTAACCGCACTTGCTTTAGG	
cad15	2614	TATTGCACTTGCTTCAGGCTATGCTTCCGGCTGAAGAAAAATTGCCTTCATTAAATGCaGGT	68/82
minnad15	2873	TATTGCACTTGCrTCAGGCTATGCCTTCAGGCTGAAGAAAAATTGCCTTCATTAAATGCGGGT	
eagand15	2913	TATTGCACTTGCTTCAGGCTATGCCTTCAGGCTGAAGAAAAATTGCCTTCATTAAATGC ACT	
pakd15	2917	TtTTGGCACTTGCTTCAGGCTATGCCTTCAGGCTGAAGAAAAATTGCCTTCATTAAATGC AGG	
sb33d15	2910	TATTGCACTTGCTTCAGGCTATGCCTTCAGGCTGAAGAAAAATTGCCTTCATTAAATGC AGG	
consensus		TATTGCACTTGCTTCAGGCTATGCCTTCAGGCTGAAGAAAAATTGCCTTCATTAAATGC -agt	

FIG. 1F. (CONTINUED)

cad15	2675	atattTTtcaAcatCacccagatcgccaaaggtagcagataaactgtatgtttaa
minnad15	2934	TATANTTTnCAAggCnagg
eagand15	2973	TATATTTTCAA
pakd15	2977	TTATATTtCAA
sb33d15	2970	TTATA

69/82

minnad15 2954

eagand15 2985

pakd15 2990

sb33d15 2975

consensus accttgttagaaatttagcaggaaaaaggatgtatggaaaatgtgtgtgt

## FIG.1F.(CONTINUED)

cad15 2797 cgtaaaaaagttagaaggccaaatggcttttagaaaaagatgcacccgcgttacgtcaag  
minnad15 2954

eagand15 2985

pakd15 2990

sb33d15 2975

consensus cgtaaaaaagttagaaggccaaatggcttttagaaaaagatgcacccgcgttacgtcaag  
70/82

cad15 2858 ctgatattcaaaaacggaggattaaataataggtgtggctgaagatgctgaatt

minnad15 2954

eagand15 2985

pakd15 2990

sb33d15 2975

consensus ctgatattcaaaaacggaggattaaataataggtgtggctgaagatgctgaatt

71/82

## FIG. 1F. (CONTINUED)

cad15 2919 acaaaaattaaatgcaagaacaaggataaaaaa

minnad15 2954

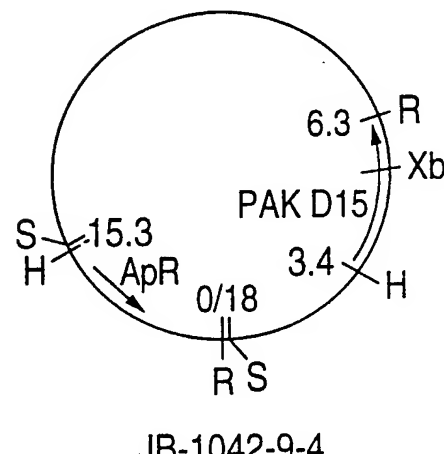
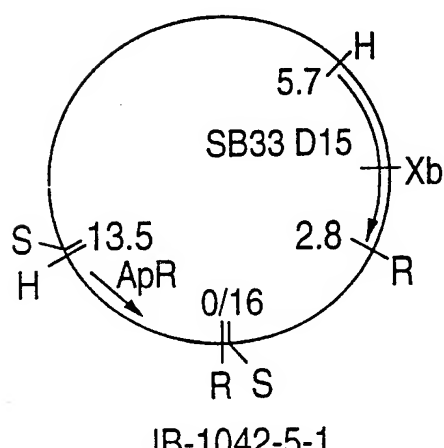
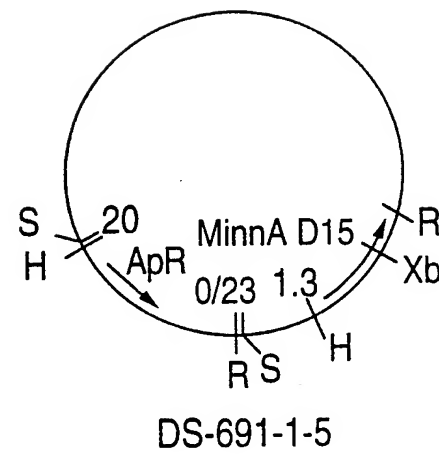
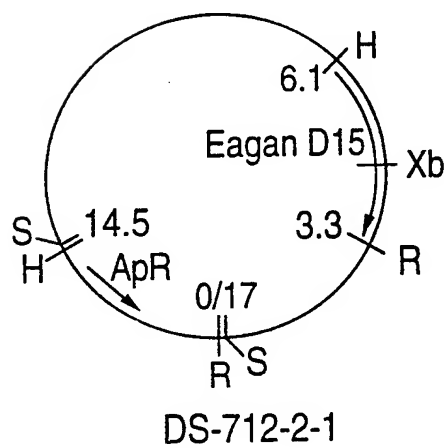
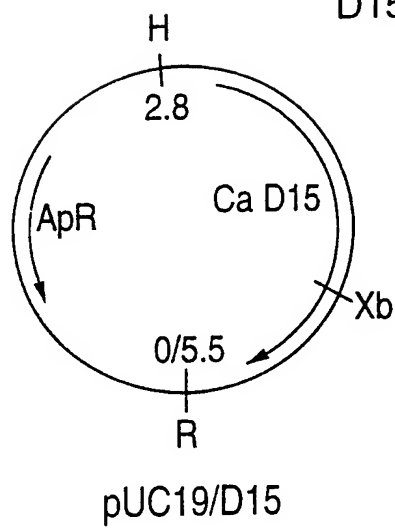
eagand15 2985

pakd15 2990

sb33d15 2975

consensus acaaaaattaaatgcaagaacaaggataaaaaa

72/82  
D15 CLONES



**FIG. 2.**

## D15 SEQUENCE COMPARISON

MKKLLIASLIFGTTTVFAAPPVAKDIRVLDGQCDLQQTRASLIPVRAGQRVITNDVANIVRSLFVSGRFDDVKAHQEQEDVILWSVVAKSIIISDVKTKGN	Ca Eagan MinnA SB33 PAK	73/82	Ca Eagan MinnA SB33 PAK	Ca Eagan MinnA SB33 PAK	Ca Eagan MinnA SB33 PAK
SVIPIEALKQNLDANGEFKVGDVLLIREKLNEPAKSVEHYASTERYNATIVEPIVTNTLPNRAELLIQIQINEDDKAKLASLTFKGNESVSSSTLQEQQMELQPD	I I				
SMWKLWGNKFFGAEQFEKDLOQSIRDYIYINGZAKAQITKTIVQINDEKIKVKNVTIDVNCEGLQYDLRSARIIGNLGMSAELEPLISAIHINDTFRSDIAD	A A				
VENAIKAKLGERRYGSATVNSVPDFDDANKTILAITLWVDAAGRILTVQLRFFENTVSADSTLROEMRQEGITWINSQQLVELGKIRLDRIGFFETVENRID	NT F F NT				

PINGSNDEVLWYKVKEKRTNGSINFGIGXGTESGSISYQASVKQDNELIGAAMSAGTKNDYGTISVNLGYTEPYFTKDGVSLGAVNFENYDKSDTSS

Eagan  
MinnA  
SB33  
PAK  
I  
T I

IVYKRITYGSNVTLGFPVNENNNSYYCELGHTYNKISNEALEYNRNLVYIQSMSKFKENGTKNDDFDSFGWYNSINRGYFPIKGVKASLGGRTTIPGSINKY

YKLSADWQGFYPLDRDHLMVSAKASAGYANGFENKRLPFYQTYTAGGIGSRLFAYGSIGPNAIYAEGNGGTCTFKLSSDVIGNAIAATASAELIV

FIG.3B.

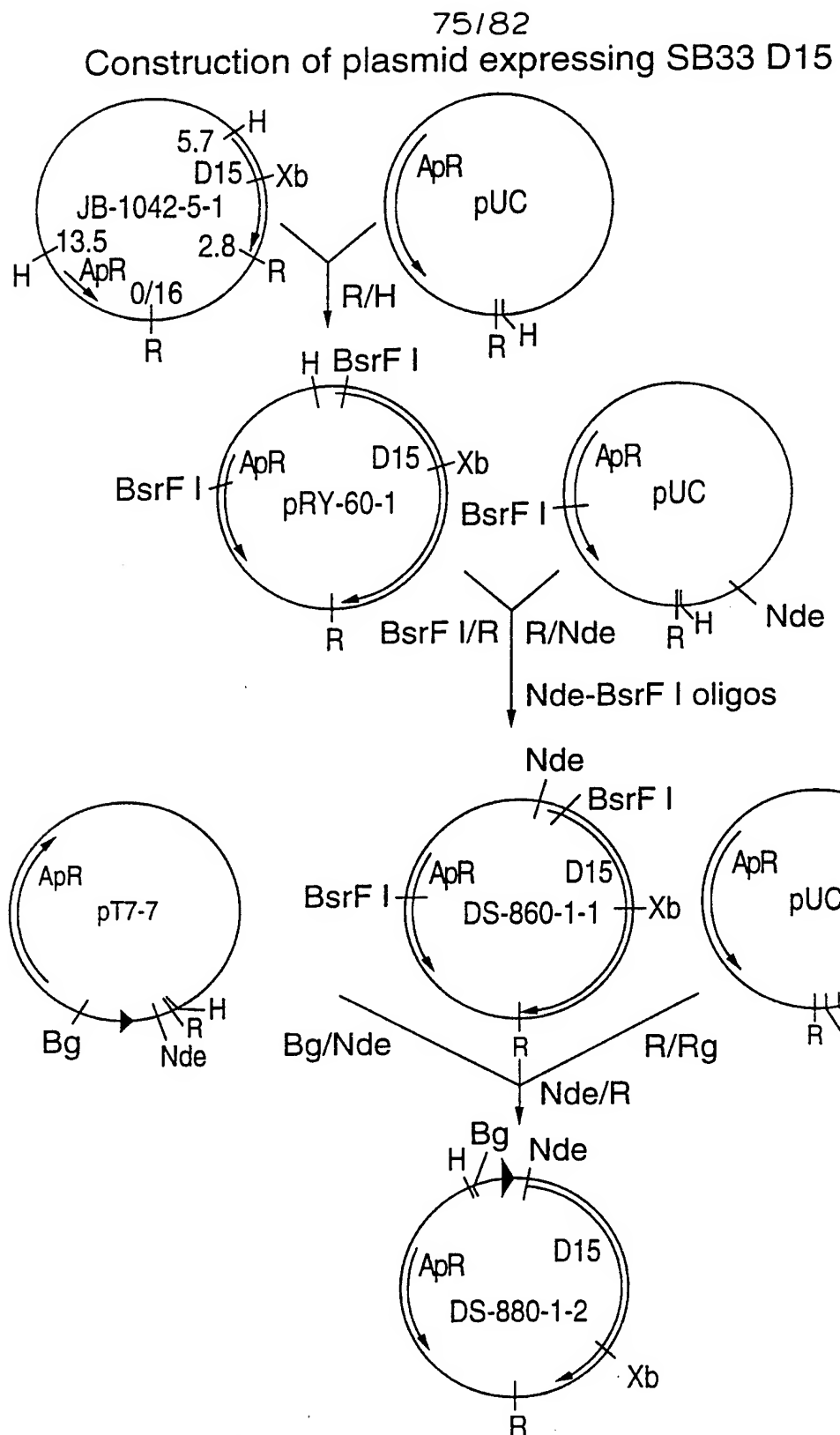
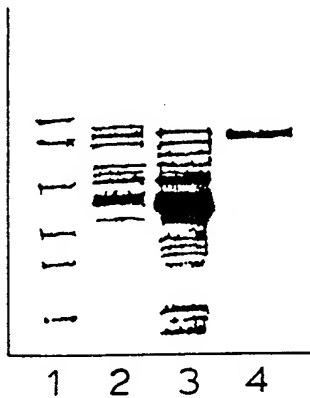


FIG.4.

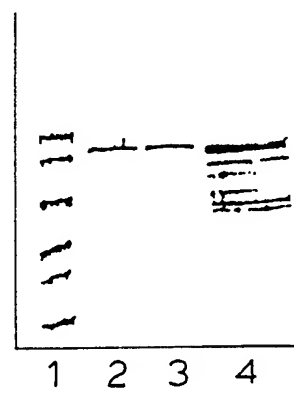
76/82

PURIFICATION OF D15 FROM A NON-TYPEABLE  
HAEMOPHILUS INFLUENZAE STRAIN 30

A



B



PROTEIN STAIN

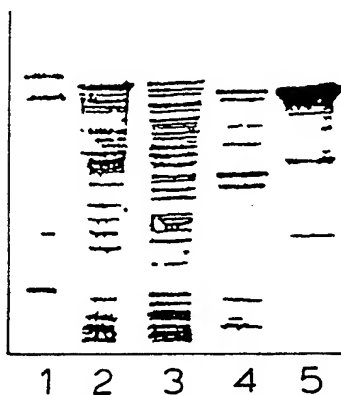
WESTERN BLOT

1. Low MW markers
2. Strain 30
3. Native D15 crude extract
4. D15 after anti-D15 affinity chromatography

FIG.5.

77 / 82

## PURIFICATION OF FULL LENGTH RECOMBINANT D15



1. Protein M.W. Markers
2. Lysate of E. coli expressed rD15
3. Soluble protein in Tris-HCl buffer extract
4. Soluble proteins in Tris/Triton X-100/ EDTA extraction buffer
5. rD15 inclusion bodies

FIG.6.

78/82

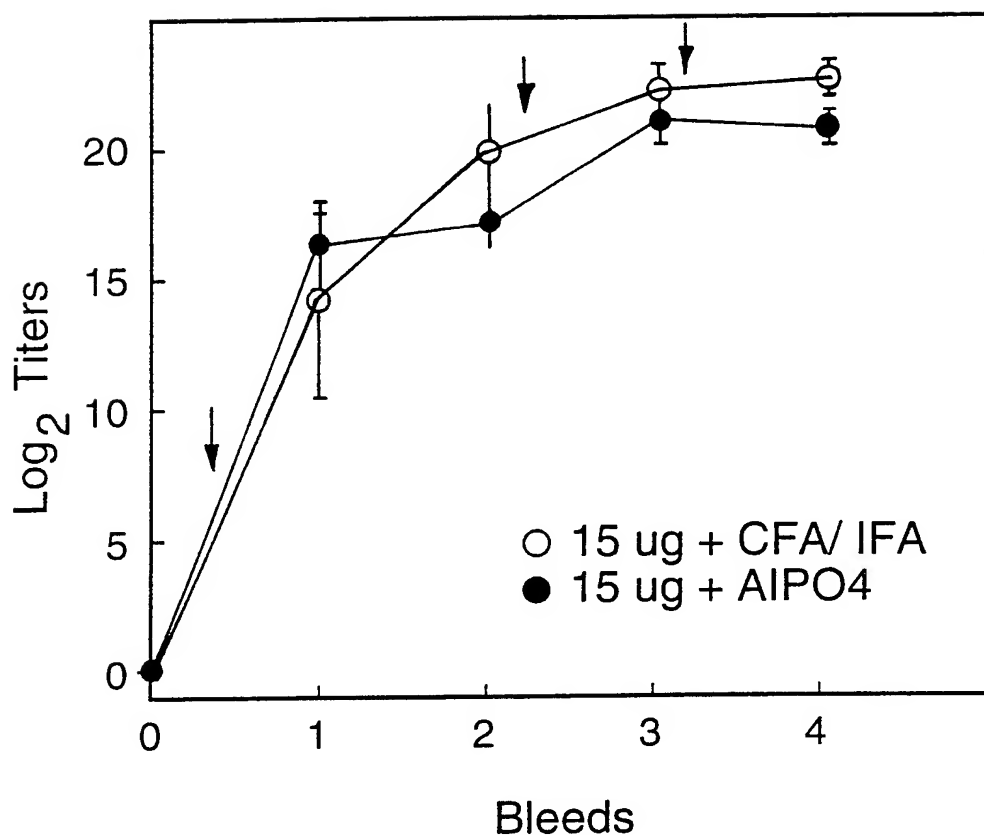


FIG. 7.

79/82

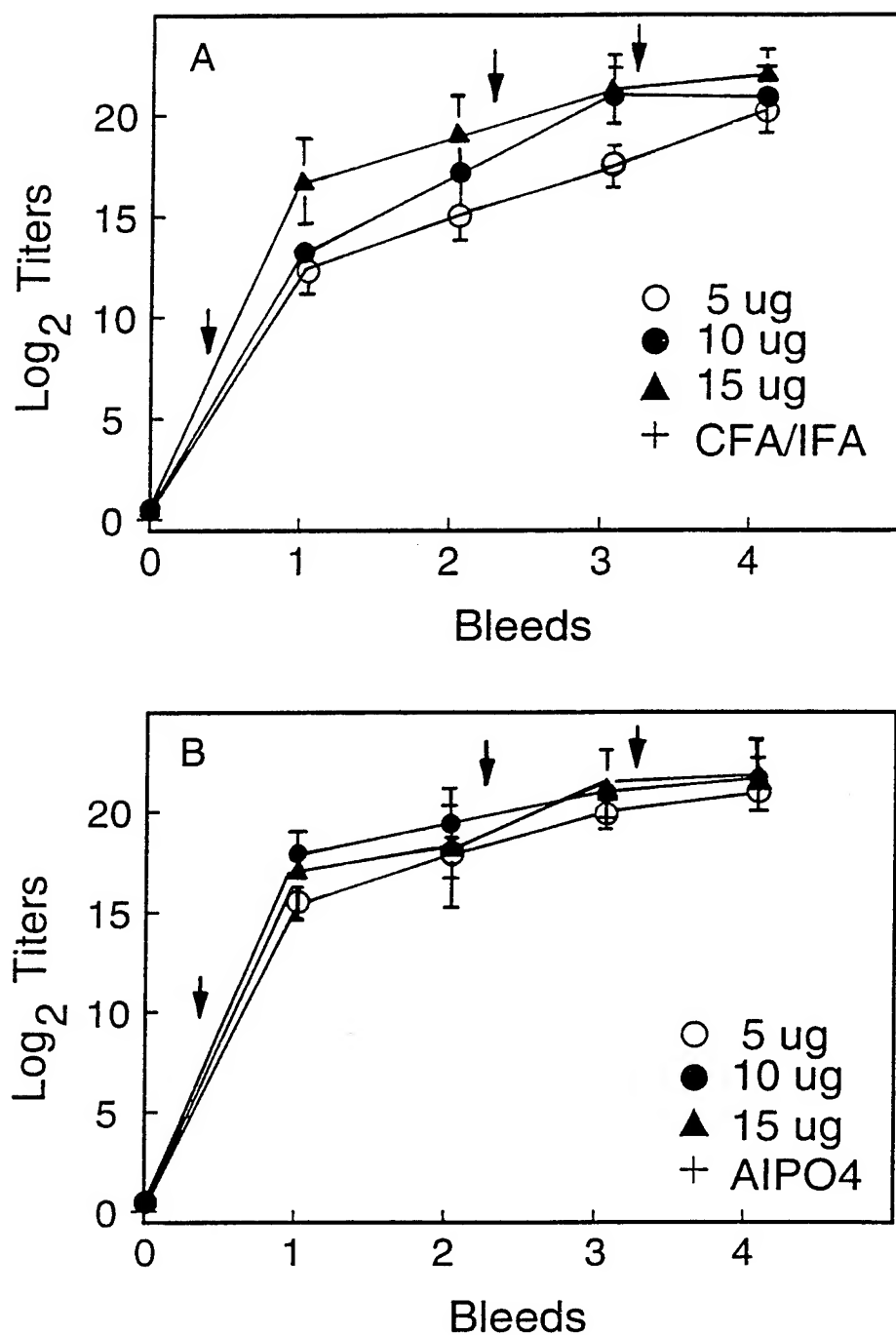
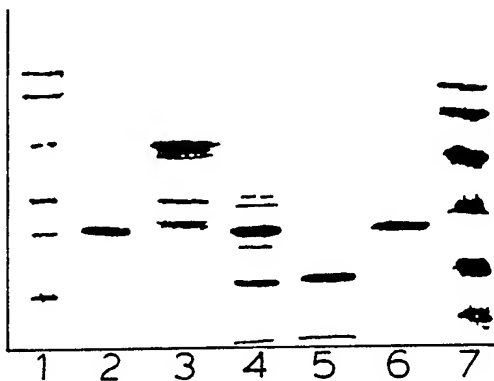


FIG.8.

80/82

PURIFICATION OF TRUNCATED D15 FROM D15-GST  
FUSION PROTEIN



1. Prestain low MW markers
2. GST standard
3. GST-(D15 fragment) fusion protein
4. Fusion protein cleaved by thrombin
5. rD15 fragment
6. GST
7. Low MW markers

FIG.9.

81/82

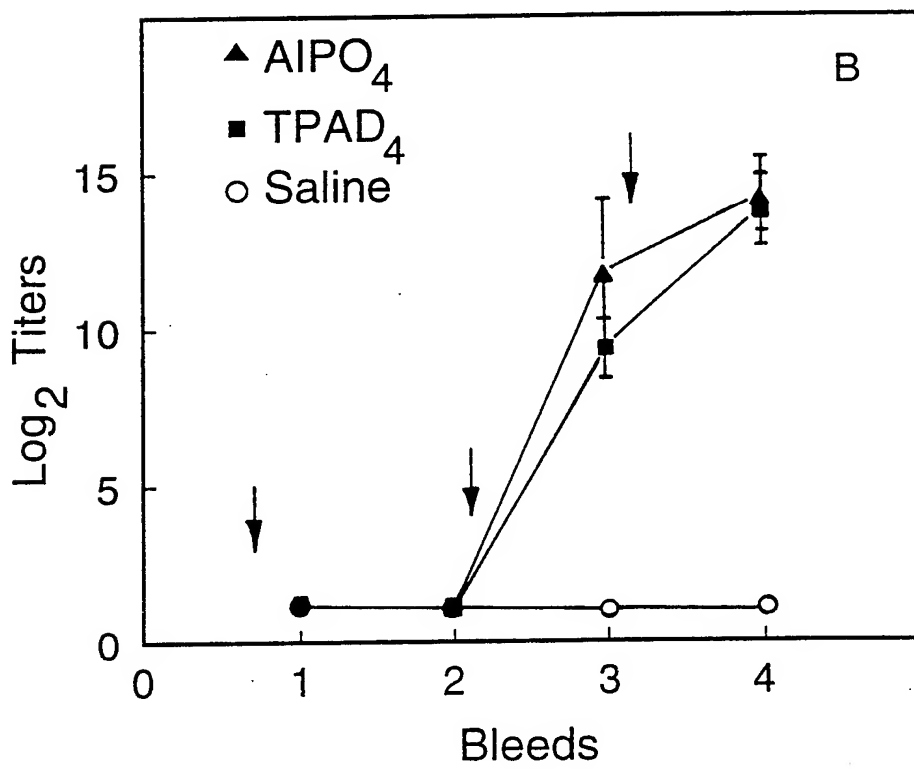
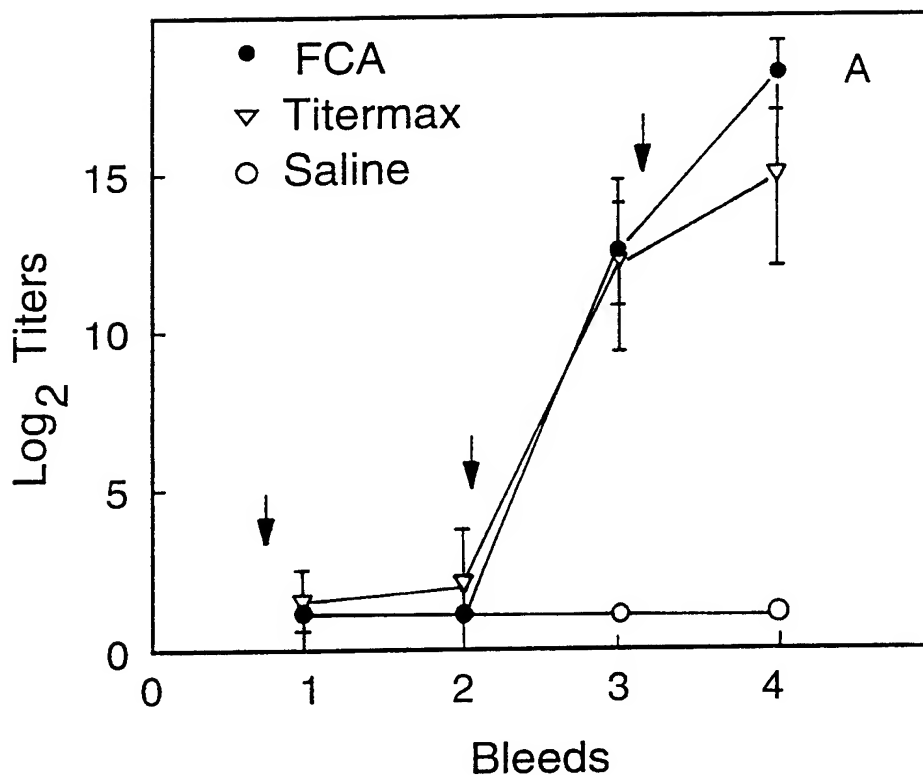
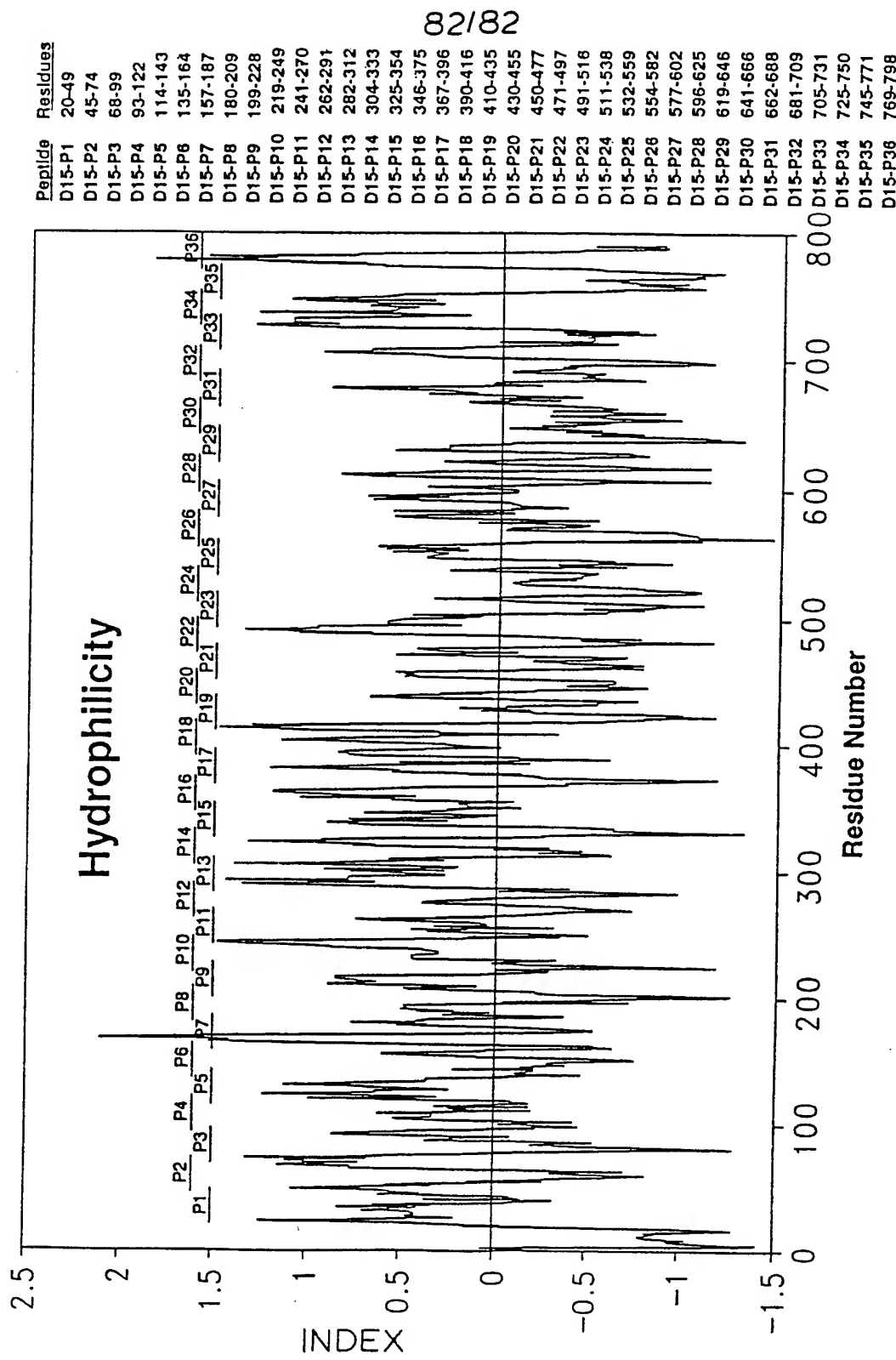


FIG.10.

FIGURE 11



## INTERNATIONAL SEARCH REPORT

In:      International Application No  
PCT/CA 93/00501

## A. CLASSIFICATION OF SUBJECT MATTER

C 12 N 15/31, C 07 K 13/00, A 61 K 39/102,  
//(C 12 N 15/31; C 12 R 1/21)

According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

A 61 K, C 07 K, C 12 N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	INFECTIOON AND IMMUNITY, vol. 58, no. 4, April 1990 W.R. THOMAS et al. "Expression in Escherichia coli of a High-Molecular-Weight Protective Surface Antigen Found in Nontypeable and Type b Haemophilus influenzae" pages 1909-1913, the whole document. --	1, 4, 6, 27
A	EP, A2, 0 378 929 (CONNAUGHT LABORATORIES LIMITED) 25 July 1990 (25.07.90), claims. --	1, 4, 6, 12, 16, 20, 22
A	WO, A1, 91/06 652	1, 4, 6,

Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

## \* Special categories of cited documents :

- \*A\* document defining the general state of the art which is not considered to be of particular relevance
- \*E\* earlier document but published on or after the international filing date
- \*L\* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- \*O\* document referring to an oral disclosure, use, exhibition or other means
- \*P\* document published prior to the international filing date but later than the priority date claimed

\*T\* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

\*X\* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

\*Y\* document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

\*&\* document member of the same patent family

Date of the actual completion of the international search

15 February 1994

Date of mailing of the international search report

25 -03- 1994

Name and mailing address of the ISA

European Patent Office, P.B. 5818 Patentlaan 2  
NL - 2280 HV Rijswijk  
Tel. (+ 31-70) 340-2040, Tx. 31 651 epo nl,  
Fax: (+ 31-70) 340-3016

Authorized officer

WOLF e.h.

## INTERNATIONAL SEARCH REPORT

-2-

Inte	onal Application No
PCT/CA 93/00501	

## C(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	(CONNAUGHT LABORATORIES LIMITED et al.) 16 May 1991 (16.05.91), claims. --	12,16, 20,22
A	EP, A1, 0 281 673 (THE RESEARCH FOUNDATION OF STATE UNIVERSITY OF NEW YORK) 14 September 1988 (14.09.88), claims. --	1,4,6, 12,18, 22,27
A	US, A, 5 013 664 (BRODEUR et al.) 07 May 1991 (07.05.91), abstract. -----	1,27

**ANHANG**

zum internationalen Recherchenbericht über die internationale Patentanmeldung Nr.

**ANNEX**

to the International Search Report to the International Patent Application No.

**NINXE**

au rapport de recherche international relatif à la demande de brevet international n°

PCT/CA 93/00501 SAE 82214

In diesem Anhang sind die Mitglieder der Patentfamilien der im obengenannten internationalen Recherchenbericht angeführten Patentdokumente angegeben. Diese Angaben dienen nur zur Orientierung und erfolgen ohne Gewähr.

This Annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The Office is in no way liable for these particulars which are given merely for the purpose of information.

La présente annexe indique les membres de la famille de brevets relatifs aux documents de brevets cités dans le rapport de recherche international visée ci-dessus. Les renseignements fournis sont donnés à titre indicatif et n'engagent pas la responsabilité de l'Office.

Im Recherchenbericht angeführtes Patentdokument Patent document cited in search report Document de brevet cité dans le rapport de recherche	Datum der Veröffentlichung Publication date Date de publication	Mitglied(er) der Patentfamilie Patent family member(s) Membre(s) de la famille de brevets	Datum der Veröffentlichung Publication date Date de publication
EP A2 378929	25-07-90	GB AO 8902178 CA AA 2006587 CA AA 2006587 EP A3 378929 JP A2 2283286 GB AO 8830124	22-03-89 22-06-90 23-06-90 01-08-90 20-11-90 22-02-89
WO A1 9106652	16-05-91	CA AA 2072095 EP A1 500576 GB AO 8924473 JP T2 5501202	01-05-91 02-09-92 20-12-89 11-03-93
EP A1 281673	14-09-88	US A 5173294 DK AO 6020/87 DK A 6020/87 JP A2 1157387	22-12-92 17-11-87 19-05-88 20-06-89
US A 5013664	07-05-91	keine - none - rien	